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TITLE OF THE INVENTION G PROTEIN-COUPLED RECEPTOR RESEMBLING GALANIN RECEPTORS

CROSS-REFERENCE TO RELATED APPLICATIONS

Not applicable.

STATEMENT REGARDING FEDERALLY-SPONSORED R&D Not applicable.

10 REFERENCE TO MICROFICHE APPENDIX Not applicable.

FIELD OF THE INVENTION

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This invention relates to novel human and rat DNAs encoding GPR54, a G protein-coupled receptor (GPCR) related to the galanin receptors, the proteins encoded by the DNAs, and methods of identifying selective agonists and antagonists of the proteins encoded by the DNAs.

BACKGROUND OF THE INVENTION

G-protein coupled receptors (GPCRs) are a very large class of membrane receptors that relay information from the exterior to the interior of cells. GPCRs function by interacting with a class of heterotrimeric proteins known as G-proteins. Most GPCRs function by a similar mechanism. Upon the binding of agonist, a GPCR catalyzes the dissociation of guanosine diphosphate (GDP) from the α subunit of G proteins. This allows for the binding of guanosine triphosphate (GTP) to the α subunit, resulting in the disassociation of the α subunit from the β and γ subunits. The freed α subunit then interacts with other cellular components, and in the process passes on the extracellular signal represented by the presence of the agonist. Occasionally, it is the freed β and γ subunits which transduce the agonist signal.

GPCRs possess common structural characteristics. They have seven hydrophobic domains, each about 20-30 amino acids long, linked by sequences of hydrophilic amino acids of varied length. These seven hydrophobic domains intercalate into the plasma membrane, giving rise to a protein with seven transmembrane domains, an extracellular amino terminus, and an intracellular

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carboxy terminus (Strader et al., 1994, Ann. Rev. Biochem. 63:101-132; Schertler et al., 1993, Nature 362:770-772l; Dohlman et al., 1991, Ann. Rev. Biochem. 60:653-688).

GPCRs are expressed in a wide variety of tissue types and respond to a wide range of ligands, e.g., protein hormones, biogenic amines, peptides, lipid derived messengers, etc. Given their wide range of expression and ligands, it is not surprising that GPCRs are involved in many pathological states. This has led to great interest in developing modulators of GPCR activity that can be used pharmacologically. For example, Table 1 of Stadel et al., 1997, Trends Pharmacol. Sci. 18:430-437, lists 37 different marketed drugs that act upon GPCRs. Accordingly, there is a great need to understand GPCR function and to develop agents that can be used to modulate GPCR activity.

Galanin is widely distributed in the central and peripheral nervous system. Galanin in most species is a 29 amino acid peptide with an amidated carboxyl terminus. Human galanin is unique in that it is longer, 30 amino acids, and is not amidated. There is strong conservation of the galanin sequence, with the amino terminal fifteen residues being absolutely conserved in all species. Galanin immunoreactivity and binding is abundant in the hypothalamus, the locus coeruleus, the hippocampus, and the anterior pituitary, as well as regions of the spinal cord, the pancreas, and the gastrointestinal tract.

Injection of galanin into the paraventricular nucleus (PVN) of the hypothalamus produces a dose-dependent increase in feeding in satiated rats. Although galanin can enhance carbohydrate ingestion, studies have shown that it profoundly increases fat intake. It has been suggested that galanin shifts macronutrient preference from carbohydrate to fat. The same injections of galanin that increase feeding reduce energy expenditure and inhibit insulin secretion. There is enhanced galanin expression in the hypothalamus of genetically obese rats compared with their lean littermates. Injection of peptide galanin receptor antagonists into the PVN blocks the galanin-specific induction of increased fat intake. Specific galanin antisense oligonucleotides when injected into the PVN produce a specific decrease in galanin expression associated with a decrease in fat ingestion and total caloric intake while hardly affecting either protein or carbohydrate intake. Thus galanin appears to be a potential neurochemical marker related to the behavior of fat ingestion and galanin receptors are attractive targets for the development of drugs to treat obesity and other eating disorders.

Galanin inhibits cholinergic function and impairs working memory in rats. Lesions that destroy cholinergic neurons result in deficits in spatial learning tasks. While locally administered acetylcholine (ACh) reverses some of this deficit, galanin blocks this ACh-mediated improvement. Evidence from autopsy samples from Alzheimer's disease-afflicted brains suggests an increased galinergic innervation of the nucleus basilis. Thus, if galinergic overactivity contributes to the decline in cognitive performance in Alzheimer's disease, galanin antagonists may be therapeutically useful in alleviating cognitive impairment.

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Other physiological processes in which galanin has been implicated include nociception (Verge et al., 1993, Neurosci. Lett. 149:193-197) and sexual behavior (Benelli et al., 1994, Eur. J. Pharmacol. 260:279-282).

In the rat, administration of galanin intracerebroventriclarly, subcutaneously, or intravenously increases plasma growth hormone. Infusion of human galanin into healthy subjects also increases plasma growth hormone and potently enhances the growth hormone response to growth hormone releasing hormone (GHRH).

Galanin levels are particularly high in dorsal root ganglia. Sciatic nerve resection dramatically up-regulates galanin peptide and mRNA levels. Chronic administration of galanin receptor antagonists (M35, M15) after axotomy results in a marked increase in self mutilation behavior in rats, generally considered to be a response to pain. Application of antisense oligonucleotides specific for galanin to the proximal end of a transected sciatic nerve suppressed the increase in galanin peptide levels with a parallel increase in autotomy. Galanin injected intrathecally acts synergistically with morphine to produce analgesia, this antinociceptive effect of morphine is blocked by galanin receptor antagonists. Thus, galanin agonists may have some utility in relieving neural pain.

The actions of galanin are mediated by at least three high affinity galanin receptors that are coupled by pertussis toxin sensitive G_i/G_0 proteins to inhibition of adenylate cyclase activity, closure of L-type Ca^{++} channels, and opening of ATP-sensitive K⁺ channels (Habert-Ortoli et al., 1994, Proc. Natl. Acad. Sci. USA 91:9780-9783; Howard et al., 1997, FEBS Lett. 405:285-290; Wang et al., 1997, J. Biol. Chem. 272:31949-31952; Kolakowski et al., 1998, J. Neurochem 71:2239-2251). Specific binding of 125 I-galanin (Kd approximately 1 nM) has been demonstrated in areas paralleling localization of galanin immunoreactivity: hypothalamus, ventral hippocampus, basal forebrain, spinal cord, pancreas, and

pituitary. In most tissues, the amino terminus (GAL 1-15) is sufficient for high affinity receptor binding and agonist activity.

A galanin receptor cDNA was isolated by expression cloning from a human Bowes melanoma cell line. (Habert-Ortoli, et al. 1994. Proc. Nat. Acad. Sci., USA 91: 9780-9783). This receptor, GALR1, is expressed in human fetal brain and small intestine, but little else is known of its distribution. Gal(1-16) is at least 1,000 times more active than pGAL(3-29) as an inhibitor of ¹²⁵I-porcine galanin binding to this receptor transiently expressed in COS cells. It remains to be determined whether this receptor subtype represents the hypothalamic receptor that mediates galanin specific feeding behavior.

Galanin receptors have been described in several international patent publications (WO 98/03548; WO 97/46681; WO 97/26853; WO 98/29439; WO 98/29440; WO 98/29441; WO 95/22608). European Patent Application EP 711830 also describes a galanin receptor.

It would be desirable to identify additional galanin receptors so that they can be used to further characterize this biological system and to identify galanin receptor subtype selective agonists and antagonists.

SUMMARY OF THE INVENTION

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The present invention is directed to novel human and rat DNAs that encode a G-protein coupled receptor, GPR54. The DNAs encoding GPR54 are substantially free from other nucleic acids and have the nucleotide sequences shown as SEQ.ID.NO.:1 (human GPR54) and SEQ.ID.NO.:2 (rat GPR54). Also provided are GPR54 proteins encoded by the novel DNA sequences. The GPR54 proteins are substantially free from other proteins and have the amino acid sequences shown as SEQ.ID.NO.:3 (human GPR54) and SEQ.ID.NO.:4 (rat GPR54). Methods of expressing GPR54 in recombinant systems and of identifying agonists and antagonists of GPR54 are provided.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1A-B shows the complete cDNA sequence and amino acid sequence of human GPR54. The DNA sequence shown is SEQ.ID.NO.:1. The amino acid sequence shown is SEQ.ID.NO.:3.

Figure 2A-B shows the complete cDNA sequence of rat GPR54 (SEQ.ID.NO.:2).

Figure 3 shows the complete amino acid sequence of human GPR54 (SEQ.ID.NO.:3).

Figure 4 shows the complete amino acid sequence of rat GPR54 (SEQ.ID.NO.:4).

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Figure 5A-B shows the location of the rat GPR54 open reading frame. The nucleotide sequence shown is (SEQ.ID.NO.:2). The amino acid sequence shown is (SEQ.ID.NO.:4).

Figure 6 shows the results of a Northern blot of rat GPR54 mRNA in rat brain. Each lane contained 5 µg of poly(A)[†] RNA isolated from various tissues.

Figure 7A-D shows darkfield autoradiograms of sagittal and coronal sections of rat brain showing the localization of GPR54 receptor mRNA. Figure 7A shows a lateral representative section at 0.9 mm. Also shown are representative sections at levels relative to the bregma at -3.3 mm (Figure 7B), -3.8 mm (Figure 7C), and -6.3 mm (Figure 7D). Aco = cortical nucleus of the amygdala; Ahy = anterior hypothalamic area; Arc = hypothalamic arcuate nucleus; IC = inferior colliculus; CA, field of Ammon's horn; DG, dentate gyrus; DM, dorsomedial hypothalamic nucleus; LC, locus coeruleus; LH, lateral hypothalamic area, LHb, lateral habenular nucleus; MeA, medial nucleus of the amygdala; MPO, medial preoptic area; MRN, mesencephalic reticular nucleus; PAG, periaqueductal gray; PB, parabrachial nucleus; PF, parafascicular thalamic nucleus; PH, posterior hypothalamic nucleus; PMV, ventral premammillary nucleus; PO, primary olfactory cortex; RSpl, retrosplenial cortex; SC, superior colliculus; SHy, septohypothalamic nucleus; VTA, ventral tegmental area; ZI, zona incerta.

Figure 8 shows an alignment of the amino acid sequence of rat GPR54 (SEQ.ID.NO.:4) with the amino acid sequence of rat GALR1 (SEQ.ID.NO.:5), rat GALR2 (SEQ.ID.NO.:6), rat GALR3 (SEQ.ID.NO.:7), and the rat opiod receptor DOR (SEQ.ID.NO.:8).

Figure 9 shows an alignment of the amino acid sequences of rat GPR54 (SEQ.ID.NO.:4) and human GPR54 (SEQ.ID.NO.:3).

DETAILED DESCRIPTION OF THE INVENTION

For the purposes of this invention:

"Substantially free from other proteins" means at least 90%, preferably 95%, more preferably 99%, and even more preferably 99.9%, free of other proteins.

Thus, a GPR54 protein preparation that is substantially free from other proteins will

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contain, as a percent of its total protein, no more than 10%, preferably no more than 5%, more preferably no more than 1%, and even more preferably no more than 0.1%, of non-GPR54 proteins. Whether a given GPR54 protein preparation is substantially free from other proteins can be determined by such conventional techniques of assessing protein purity as, e.g., sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) combined with appropriate detection methods, e.g., silver staining or immunoblotting.

"Substantially free from other nucleic acids" means at least 90%, preferably 95%, more preferably 99%, and even more preferably 99.9%, free of other nucleic acids. Thus, a GPR54 DNA preparation that is substantially free from other nucleic acids will contain, as a percent of its total nucleic acid, no more than 10%, preferably no more than 5%, more preferably no more than 1%, and even more preferably no more than 0.1%, of non-GPR54 nucleic acids. Whether a given GPR54 DNA preparation is substantially free from other nucleic acids can be determined by such conventional techniques of assessing nucleic acid purity as, e.g., agarose gel electrophoresis combined with appropriate staining methods, e.g., ethidium bromide staining, or by sequencing.

"Functional equivalent" means a receptor which does not have exactly the same amino acid sequence as naturally occurring GPR54, due to alternative splicing, substitutions, deletions, mutations, or additions, but retains substantially the same biological activity as GPR54. Such functional equivalents will have significant amino acid sequence identity with naturally occurring GPR54. Genes and DNA encoding such functional equivalents can be detected by reduced stringency hybridization with a DNA sequence encoding naturally occurring GPR54. For the purposes of this invention, naturally occurring GPR54 has the amino acid shown as SEQ.ID.NO.:3 or SEQ.ID.NO.:4. A nucleic acid encoding a functional equivalent has at least about 50% identity at the nucleotide sequence level to SEQ.ID.NO.:1 or SEQ.ID.NO.:2.

A polypeptide has "substantially the same biological activity" as GPR54 if that polypeptide has a K_d for a ligand that is no more than 5-fold greater than the K_d of GPR54 having SEQ.ID.NO.:3 or SEQ.ID.NO.:4 for the same ligand. A polypeptide also has "substantially the same biological activity" as GPR54 if that polypeptide is capable of mediating the same functional response as naturally occurring GPR54 when exposed to the same ligand as naturally occurring GPR54. Examples of functional responses are: pigment aggregation in *Xenopus* melanophores,

changes in membrane currents in Xenopus oocytes, modulation of cAMP levels, changes in calcium concentration, changes in inositol phosphate levels, and coupling to inwardly rectifying potassium channels. One skilled in the art would be familiar with a variety of methods of measuring the functional responses of other G-protein coupled receptors and would be able to apply those methods to GPR54 (see, e.g., 5 Lerner, 1994, Trends Neurosci. 17:142-146 [changes in pigment distribution in melanophore cells]; Yokomizo et al., 1997, Nature 387:620-624 [changes in cAMP or calcium concentration, chemotaxis]; Howard et al., 1996, Science 273:974-977 [changes in membrane currents in Xenopus oocytes]; McKee et al., 1997, Mol. Endocrinol. 11:415-423 [changes in calcium concentration measured using the 10 aequorin assay]; Offermanns & Simon, 1995, J. Biol. Chem. 270:15175, 15180 [changes in inositol phosphate levels]). Zlokarnik et al., 1998, Science 279:84-88 and U.S. Patent No. 5,741,657 describe a reporter gene assay that can be adapted to measure GPR54 functional responses. The assay utilizes an inducible promoterdriven \u03b3-lactamase that cleaves a fluorescent substrate. Cleavage of the substrate 15 leads to a change in fluoresence resonance energy transfer (FRET) between different portions of the substrate that is proportional to the magnitude of induction of the βlactamase. Thus, the level of activation of the inducible promoter determines the amount of FRET measured. This level of induction of the promoter is in turn determined by the level of the substance (e.g., cAMP) the promoter is induced by. By 20 choosing a promoter that is induced by a functional response that results from the interaction of a ligand and GPR54 (e.g., changes in cAMP levels), one can use this assay to measure GPR54 functional responses.

Depending upon the cells in which GPR54 is expressed, and thus the G-proteins with which GPR54 is coupled, certain of such methods as described above may be appropriate for measuring the functional responses of GPR54. It is well within the competence of one skilled in the art to select the appropriate method of measuring functional responses for a given experimental system.

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A "conservative amino acid substitution" refers to the replacement of one amino acid residue by another, chemically similar, amino acid residue. Examples of such conservative substitutions are: substitution of one hydrophobic residue (isoleucine, leucine, valine, or methionine) for another; substitution of one polar residue for another polar residue of the same charge (e.g., arginine for lysine; glutamic acid for aspartic acid).

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By "isolated GPR54 protein" or "isolated GPR54 DNA" is meant GPR54 protein or DNA encoding GPR54 that has been isolated from a natural source. Use of the term "isolated" indicates that GPR54 protein or DNA has been removed from its normal cellular environment. Thus, an isolated GPR54 protein may be in a cell-free solution or placed in a different cellular environment from that in which it occurs naturally. The term isolated does not imply that an isolated GPR54 protein is the only protein present, but instead means that an isolated GPR54 protein is at least 95% free of non-amino acid material (e.g., nucleic acids, lipids, carbohydrates) naturally associated with the GPR54 protein. Thus, a GPR54 protein that is expressed in bacteria or even in eukaryotic cells which do not naturally (i.e., without human intervention) express it through recombinant means is an "isolated GPR54 protein." Similarly, DNA encoding GPR54 that is present in bacteria or even in eukaryotic cells which do not naturally (i.e., without human intervention) contain it through recombinant means is an "isolated DNA encoding GPR54."

The present invention pertains to the discovery of DNA encoding a galanin receptor-like protein. Two degenerate primers (P1 and P2, see Example 1) based on conserved GPCR sequences in transmembrane segment 3 (TM3) and transmembrane segment 7 (TM7), respectively, were used to amplify an aliquot of a rat brain cDNA library with proof-reading Pfu polymerase. The amplified DNA was excised and subcloned into the pBluescript vector. One of the resulting rat clones appeared to partially encode a galanin/opioid-like receptor. The partial cDNA was labeled with ^{32}P dCTP- α and used to screen the cDNA library employed in the degenerate PCR. Two positive plaques were purified and their inserts amplified by PCR using Pfu polymerase and primers flanking the cloning site of the λ gt11 vector. The PCR products were subcloned into pBluescript and sequenced. Sequence analysis revealed that each plaque encoded a region of a putative GPCR from TM3 to

analysis revealed that each plaque encoded a region of a putative GPCR from TM3 to the carboxy terminus identical to each other and the original probe. A second round of screening of 1×10^6 plaques freshly plated from the same library yielded an additional three positive plaques. PCR amplification of these positive plaques with $\lambda gt11$ flanking primers, each paired with an internal primer, revealed that only one of these positive plaques contained the entire open reading frame (ORF). This plaque was purified, the insert subcloned into pBluescript and was confirmed to contain the 5' end of the full-length open reading frame. Finally, two specific primers from the 5' and 3' ends of the ORF were used to amplify with pfu polymerase the full length rat

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cDNA 1.2 Kb clone, named GPR54. Sequence analysis revealed the cloned GPR54 ORF to be identical to the previous phage clones and the original probe.

GPR54 contained an ORF of 1,185 bp encoding a protein of 395 amino acids.

Using GPR54 in a BLAST search (Altschul, 1997, Nucleic Acids Res 25:3389-3402), the highest identity was observed with the galanin and opioid receptor families. Specifically, GPR54 shared an amino acid sequence identity in the TM regions with rat galanin receptors GalR1(45%), GalR3 (45%), GalR2 (44%), and rat opioid receptor DOR (37%) (Figure 8). Conserved residues and consensus sequences of the rhodopsin superfamily of GPCRs present in GPR54 included an asparagine in TM1, an aspartate in TM2, prolines in TMs 4 through 7, three consensus sequences for N-linked glycosylation in the amino terminus, cysteines in the first and second extracellular loops, a PKA/PKC consensus sequence in the second intracellular loop. a PKC consensus sequence in the third intracellular loop, and three possible palmitoylation cysteine sites in the carboxy tail. Significantly, various residues in the human GalR1 receptor shown to be important for high-affinity galanin binding (corresponding to His262, His265, Glu269, and Phe280 in rat GalR1; (Kask et al., 1996, EMBO J. 15:236-244 (Kask); Berthold et al., 1997, Eur. J. Biochem. 249:601-606 (Berthold)) were not conserved in GPR54. Among these however, only His262 is conserved among the three galanin receptors. In addition, the substitution of a tyrosine residue found in GPR54, GalR2 and GalR3 in place of Phe280 in GalR1 was shown to have no significant effect on galanin binding (Kask) as opposed to previous studies where Phe280 was replaced by alanine in GalR1 (Berthold).

Both Northern blot and *in situ* hybridization analyses of GPR54 were performed at high stringencies and with a DNA probe encoding GPR54 from TM3 to TM7 and with low identities to the genes encoding galanin and related receptors. The tissue distribution of GPR54 was obtained by northern blot analysis using poly(A)[†] RNA isolated from various rat tissues (Figure 6). In the brain, multiple RNA transcripts with a complex pattern were detected in the medulla pons, midbrain, hippocampus, cortex, frontal cortex, and striatum. The most intense band was approximately 3.7 Kb in length, with a single, larger transcript of approximately 12 Kb length detected in the liver and intestine only. No transcripts were revealed in the cerebellum or kidney tissues.

Using *in situ* hybridization of rat brain sections, the distribution of GPR54 mRNA was found to be discretely localized to many areas (Figure 7). The

highest levels of expression were seen in hypothalamic and amygdaloid nuclei. GPR54 mRNA was highly expressed in the zona incerta, ventral tegmental area, dentate gyrus, hypothalamic arcuate nucleus, dorsomedial hypothalamic nucleus, primary olfactory cortex, lateral habenular nucleus, lateral hypothalamic area, locus coeruleus, and the cortical and medial nuclei of the amygdala. GPR54 mRNA was also concentrated in the superior colliculus, medial preoptic area, anterior hypothalamic area, posterior hypothalamic nucleus, periaqueductal gray, parafascicular thalamic nucleus, parabrachial nucleus, and ventral premammillary nucleus. The signals detected in the septohypothalamic nucleus, inferior colliculus, medial nucleus of the amygdala, mesencephalic reticular nucleus and retrosplenial cortex were diffuse and less abundant.

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GPR54's CNS expression pattern was found to resemble those of galanin receptors. Specifically, rat GalR1 mRNA expression is abundant in several brain regions including the hypothalamus, amygdala, hippocampus and locus coeruleus (Parker et al., 1995, Mol. Brain Res. 34:179-189). Rat GalR2 mRNA expression is found in the mammilary nuclei, the dentate gyrus and posterior hypothalamic and arcuate nuclei (Kolakowski et al., 1998, J. Neurochem. 71, 2239-2251). Rat GalR3 is found to be abundantly expressed in the CA regions of Ammon's horn and the dentate gyrus with transcripts also detected in thalamic, hypothalamic, mammilary and amygdaloid nuclei (Kolakowski et al., 1998, J. Neurochem. 71, 2239-2251).

The identity and overlapping expression patterns of GPR54 with the galanin receptors suggested that the encoded receptor may demonstrate binding to galanin. In preparation for expression and binding studies, the 1.2 kb cDNA fragment encoding the ORF of GPR54 was subcloned into the multiple cloning site of the pcDNA3 expression vector and transiently transfected into COS-7 cells. No specific binding was observed with ¹²⁵I-human galanin. In contrast, specific and high affinity binding was observed under similar conditions with ¹²⁵I-human galanin in membranes prepared from COS cells transfected with human GalR1, consistent with a previous report for GalR2 and GalR3 (Kolakowski et al., 1998, J. Neurochem. 71, 2239-2251).

A BLAST search with the rat GPR54 sequence revealed high identity with a human 3.5 Mb contig located in chromosome 19p13.3 containing a serine protease gene cluster (GenBank accession number AC005379). Sequence analysis revealed a previously unrecognized 3.3 kb intron-containing human orthologue of

GPR54 encoding a protein 398 amino acids in length and sharing a translated amino acid identity of 81% (100% identity in the TM regions) with rat GPR54. The genomic sequence revealed four introns located in TM2 (~800 bp, interrupting the translated FYI..ANL sequence), TM3 (~800 bp, interrupting IQQ..VSV), TM4 (~250 bp, interrupting WVG..SAA) and in the third intracellular loop (~180 bp, interrupting ALQ..GQV).

One aspect of this invention is an isolated DNA comprising nucleotides encoding a polypeptide having the amino acid sequence SEQ.ID.NO.:3 or SEQ.ID.NO.:4. This isolated DNA can be substantially free from other nucleic acids and can be either single stranded or double stranded, *i.e.*, paired with its complementary sequence. Also within the present invention is isolated RNA corresponding to this DNA.

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Another aspect of this invention is the identification and cloning of a cDNA which encodes GPR54, a G protein-coupled receptor. This cDNA is substantially free from other nucleic acids and can be either single stranded or double stranded. The present invention provides a cDNA molecule substantially free from other nucleic acids having the nucleotide sequence shown in Figure 1 as SEQ.ID.NO.:1 or in Figure 2 as SEQ.ID.NO.:2. SEQ.ID.NO.:1 contains an open reading frame (positions 1-1,194 of SEQ.ID.NO.:1) encoding a protein of 398 amino acids. SEQ.ID.NO.:2 contains an open reading frame (positions 61-1,245 of SEQ.ID.NO.:2) encoding a protein of 395 amino acids. (see Figure 5A-B).

Thus, the present invention also provides a DNA molecule substantially free from other nucleic acids comprising the nucleotide sequence of positions 1-1,194 of SEQ.ID.NO.:1 as well as a DNA molecule substantially free from other nucleic acids comprising the nucleotide sequence of positions 61-1,245 of SEQ.ID.NO.:2. The present invention also provides recombinant DNA molecules comprising the nucleotide sequence of positions 1-1,194 of SEQ.ID.NO.:1 or positions 61-1,245 of SEQ.ID.NO.:2.

Based on their predicted amino acid sequences, the human and rat

30 GPR54 proteins most likely represent novel G-protein coupled receptors (GPCRs)

since these GPR54 proteins obtain many of the characteristic features of GPCRs, e.g..

- (a) seven transmembrane domains;
- (b) three intracellular loops;
- (c) three extracellular loops; and
- (d) the GPCR triplet signature sequence.

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Northern blot and *in situ* hybridization analyses such as Figure 6 and Figure 7 showed that GPR54 RNA is widely expressed in rat brain regions (pons, midbrain, thalamus, hypothalamus, hippocampus, amygdala, cortex, frontal cortex, and striatum) as well as peripheral regions (liver and intestine).

The novel DNA sequences of the present invention encoding GPR54, in whole or in part, can be linked with other DNA sequences, *i.e.*, DNA sequences to which GPR54 is not naturally linked, to form "recombinant DNA molecules" containing GPR54 sequences. The novel DNA sequences of the present invention can be inserted into vectors in order to direct recombinant expression of GPR54. Such vectors may be comprised of DNA or RNA; for most purposes DNA vectors are preferred. Typical vectors include plasmids, modified viruses, bacteriophage, cosmids, yeast artificial chromosomes, and other forms of episomal or integrated DNA that can encode GPR54. One skilled in the art can readily determine an appropriate vector for a particular use.

Included in the present invention are DNA sequences that hybridize to SEQ.ID.NO.:1 or SEQ.ID.NO.:2 under stringent conditions. By way of example, and not limitation, a procedure using conditions of high stringency is as follows: Prehybridization of filters containing DNA is carried out for 2 hr. to overnight at 65°C in buffer composed of 6X SSC, 5X Denhardt's solution, and 100 µg/ml denatured salmon sperm DNA. Filters are hybridized for 12 to 48 hrs at 65°C in prehybridization mixture containing 100 µg/ml denatured salmon sperm DNA and 5-20 X 10⁶ cpm of 32P-labeled probe. Washing of filters is done at 37°C for 1 hr in a solution containing 2X SSC, 0.1% SDS. This is followed by a wash in 0.1X SSC, 0.1% SDS at 50°C for 45 min. before autoradiography.

Other procedures using conditions of high stringency would include either a hybridization step carried out in 5XSSC, 5X Denhardt's solution, 50% formamide at 42°C for 12 to 48 hours or a washing step carried out in 0.2X SSPE, 0.2% SDS at 65°C for 30 to 60 minutes.

Reagents mentioned in the foregoing procedures for carrying out high stringency hybridization are well known in the art. Details of the composition of these reagents can be found in, e.g., Sambrook, Fritsch, and Maniatis, 1989, Molecular Cloning: A Laboratory Manual, second edition, Cold Spring Harbor Laboratory Press. In addition to the foregoing, other conditions of high stringency which may be used are well known in the art.

The degeneracy of the genetic code is such that, for all but two amino acids, more than a single codon encodes a particular amino acid. This allows for the construction of synthetic DNA that encodes the GPR54 protein where the nucleotide sequence of the synthetic DNA differs significantly from the nucleotide sequence of SEQ.ID.NO.:1 or SEQ.ID.NO.:2, but still encodes the same GPR54 protein as SEQ.ID.NO.:1 or SEQ.ID.NO.:2. Such synthetic DNAs are intended to be within the scope of the present invention. If it is desired to express such synthetic DNAs in a particular host cell or organism, the codon usage of such synthetic DNAs can be adjusted to reflect the codon usage of that particular host cell or organism, thus leading to higher levels of expression of GPR54 protein in the host.

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Another aspect of the present invention includes host cells that have been engineered to contain and/or express DNA sequences encoding GPR54. Such recombinant host cells can be cultured under suitable conditions to produce GPR54. An expression vector containing DNA encoding GPR54 can be used for expression of GPR54 in a recombinant host cell. Recombinant host cells may be prokaryotic or eukaryotic, including but not limited to, bacteria such as E. coli, fungal cells such as yeast, mammalian cells including, but not limited to, cell lines of human, bovine, porcine, monkey, and rodent origin, and insect cells including but not limited to, Drosophila and silkworm derived cell lines. Cell lines derived from mammalian species which are suitable for recombinant expression of GPR54 and which are commercially available, include but are not limited to, L cells L-M(TK-) (ATCC CCL 1.3), L cells L-M (ATCC CCL 1.2), HEK 293 (ATCC CRL 1573), Raji (ATCC CCL 86), CV-1 (ATCC CCL 70), COS-1 (ATCC CRL 1650), COS-7 (ATCC CRL 1651), CHO-K1 (ATCC CCL 61), 3T3 (ATCC CCL 92), NIH/3T3 (ATCC CRL 1658), HeLa (ATCC CCL 2), C127I (ATCC CRL 1616), BS-C-1 (ATCC CCL 26), MRC-5 (ATCC CCL 171), Xenopus melanophores, and Xenopus oocytes.

Human embryonic kidney (HEK 293) cells and Chinese hamster ovary (CHO) cells are particularly suitable for expression of the GPR54 protein because these cells express a large number of G-proteins. Thus, it is likely that at least one of these G-proteins will be able to functionally couple the signal generated by interaction of GPR54 and its ligands, thus transmitting this signal to downstream effectors, eventually resulting in a measurable change in some assayable component, e.g., cAMP level, expression of a reporter gene, hydrolysis of inositol lipids, or intracellular Ca²⁺ levels.

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Other cells that are particularly suitable for expression of the GPR54 protein are immortalized melanophore pigment cells from *Xenopus laevis*. Such melanophore pigment cells can be used for functional assays using recombinant expression of GPR54 in a manner similar to the use of such melanophore pigment cells for the functional assay of other recombinant GPCRs (Graminski et al., 1993, J. Biol. Chem. 268:5957-5964; Lerner, 1994, Trends Neurosci. 17:142-146; Potenza & Lerner, 1992, Pigment Cell Res. 5:372-378).

A variety of mammalian expression vectors can be used to express recombinant GPR54 in mammalian and other cells. Commercially available mammalian expression vectors which are suitable include, but are not limited to, pCR2.1 (Invitrogen), pMC1neo (Stratagene), pSG5 (Stratagene), pcDNAI and pcDNAIamp, pcDNA3, pcDNA3.1, pCR3.1 (Invitrogen), EBO-pSV2-neo (ATCC 37593), pBPV-1(8-2) (ATCC 37110), pdBPV-MMTneo(342-12) (ATCC 37224), pRSVgpt (ATCC 37199), pRSVneo (ATCC 37198), and pSV2-dhfr (ATCC 37146). For expression in non-mammalian cells, various suitable expression vectors are known in the art. The choice of vector will depend upon cell type used, level of expression desired, and the like. Following expression in recombinant cells, GPR54 can be purified to a level that is substantially free from other proteins by conventional techniques, *e.g.*, salt fractionation, ion exchange chromatography, size exclusion chromatography, hydroxylapatite adsorption chromatography, hydrophobic interaction chromatography, and preparative gel electrophoresis.

The present invention includes GPR54 protein substantially free from other proteins. The amino acid sequence of the full-length human GPR54 protein is shown in Figure 3 as SEQ.ID.NO.:3. The amino acid sequence of the full-length rat GPR54 protein is shown in Figure 4 as SEQ.ID.NO.:4. Thus, the present invention includes GPR54 proteins substantially free from other proteins having the amino acid sequence of SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

As with many receptor proteins, it is possible to modify many of the amino acids of GPR54, particularly those which are not found in the ligand binding domain, and still retain substantially the same biological activity as the original receptor. Thus, the present invention includes modified GPR54 polypeptides which have amino acid deletions, additions, or substitutions but that still retain substantially the same biological activity as naturally occurring GPR54. It is generally accepted that single amino acid substitutions do not usually alter the biological activity of a protein (see, e.g., Molecular Biology of the Gene, Watson et al., 1987, Fourth Ed.,

The Benjamin/Cummings Publishing Co., Inc., page 226; and Cunningham & Wells, 1989, Science 244:1081-1085). Accordingly, the present invention includes polypeptides where one amino acid substitution has been made in SEQ.ID.NO.:3 or SEQ.ID.NO.:4 wherein the polypeptides still retain substantially the same biological activity as naturally occurring GPR54. The present invention also includes polypeptides where two or more amino acid substitutions have been made in SEQ.ID.NO.:3 or SEQ.ID.NO.:4 wherein the polypeptides still retain substantially the same biological activity as naturally occurring GPR54. In particular, the present invention includes embodiments where the above-described substitutions are conservative substitutions. In particular, the present invention includes embodiments where the above-described substitutions do not occur in the ligand-binding domain of GPR54.

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When deciding which amino acid residues of GPR54 may be substituted to produce polypeptides that are functional equivalents of GPR54, one skilled in the art would be guided by a comparison of the amino acid sequence of GPR54 with the amino acid sequences of related proteins, e.g., the human, mouse, or rat GALR1, GALR2, or GALR3 receptors, as well as the rat opiod receptor DOR (see, e.g., Figure 8). Such a comparison would allow one skilled in the art to minimize the number of amino acid substitutions made in regions that are highly conserved between GPR54 and the related proteins. Accordingly, the present invention includes polypeptides where two or more amino acid substitutions have been made in SEQ.ID.NO.:3 or SEQ.ID.NO.:4 where the polypeptides still retain substantially the same biological activity as naturally occurring GPR54 and where the substitutions are conservative and do not occur in positions where GPR54 and any of the human, mouse, or rat GALR1, GALR2, or GALR3 receptors share the same amino acid, or do not occur in positions where GPR54 and the rat opiod DOR receptor share the same amino acid (see Figure 8). In particular embodiments, the substitutions do not occur in positions where GPR54 and any of the rat GALR1, GALR2, or GALR3 receptors share the same amino acid (see Figure 8).

One skilled in the art would also recognize that polypeptides that are functional equivalents of GPR54 and have changes from the GPR54 amino acid sequence that are small deletions or insertions of amino acids could also be produced by following the same guidelines, *i.e.*, minimizing the differences in amino acid sequence between GPR54 and related proteins. Small deletions or insertions are generally in the range of about 1 to 5 amino acids. The effect of such small deletions

or insertions on the biological activity of the modified GPR54 polypeptide can easily be assayed by producing the polypeptide synthetically or by making the required changes in DNA encoding GPR54 and then expressing the DNA recombinantly and assaying the protein produced synthetically or by such recombinant expression.

Assays that could be used include simple binding assays to determine if the modified GPR54 polypeptide is capable of binding the same ligands, with approximately the same affinity, as naturally occurring GPR54 protein. Alternatively, one can use functional assays such as assays such as those described herein.

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The present invention also includes C-terminal truncated forms of GPR54, particularly those which encompass the extracellular portion of the receptor, but lack the intracellular signaling portion of the receptor. Such truncated receptors are useful in various binding assays described herein, for crystallization studies, and for structure-activity-relationship studies.

The present invention also includes chimeric GPR54 proteins.

Chimeric GPR54 proteins consist of a contiguous polypeptide sequence of GPR54 fused in frame to a polypeptide sequence of a non-GPR54 protein. For example, the N-terminal domain and seven transmembrane spanning domains of GPR54 fused at the C-terminus in frame to a G protein would be a chimeric GPR54 protein.

The present invention also includes GPR54 proteins that are in the form of multimeric structures, e.g., dimers. Such multimers of other G-protein coupled receptors are known (Hebert et al., 1996, J. Biol. Chem. 271, 16384-16392; Ng et al., 1996, Biochem. Biophys. Res. Comm. 227, 200-204; Romano et al., 1996, J. Biol. Chem. 271, 28612-28616). The dimers may be homodimers containing two GPR54 proteins or the dimers may be heterodimers containing GPR54 and another protein.

The present invention also includes isolated forms of GPR54 proteins. The present invention includes methods of identifying compounds that specifically bind to GPR54 protein, as well as compounds identified by such methods. The specificity of binding of compounds having affinity for GPR54 is shown by measuring the affinity of the compounds for recombinant cells expressing the cloned receptor or for membranes from such cells. Expression of the cloned receptor and screening for compounds that bind to GPR54, or that inhibit the binding of a known ligand of GPR54 to such cells, or membranes prepared from such cells, provides an effective method for the rapid selection of compounds with high affinity for GPR54. Such ligands or compounds can be radiolabeled, but can also be nonisotopic

compounds that can be used to displace bound radiolabeled ligands or that can be used as activators or inhibitors in functional assays. Compounds identified by the above method are likely to be agonists or antagonists of GPR54 and may be peptides, proteins, or non-proteinaceous organic molecules. Such compounds are likely to be pharmacologically useful modulators of GPR54 activity.

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Therefore, the present invention includes assays by which GPR54 agonists and antagonists may be identified. Methods for identifying agonists and antagonists of other receptors are well known in the art and can be adapted to identify agonists and antagonists of GPR54. Accordingly, the present invention includes a method for determining whether a substance is a potential agonist or antagonist of GPR54 that comprises:

- (a) transfecting cells with an expression vector encoding GPR54;
- (b) allowing the transfected cells to grow for a time sufficient to allow GPR54 to be expressed;
- (c) exposing the cells to a labeled ligand of GPR54 in the presence and in the absence of the substance;
- (d) measuring the binding of the labeled ligand to GPR54; where if the amount of binding of the labeled ligand is less in the presence of the substance than in the absence of the substance, then the substance is a potential agonist or antagonist of GPR54.

The conditions under which step (c) of the method is practiced are conditions that are typically used in the art for the study of protein-ligand interactions: e.g., physiological pH; salt conditions such as those represented by such commonly used buffers as PBS or in tissue culture media; a temperature of about 4°C to about 55°C.

The present invention also includes a method for determining whether a substance is capable of binding to GPR54, *i.e.*, whether the substance is a potential agonist or an antagonist of GPR54, where the method comprises:

- (a) providing test cells by transfecting cells with an expression vector that directs the expression of GPR54 in the cells;
 - (b) exposing the test cells to the substance;
 - (c) measuring the amount of binding of the substance to GPR54 in the test cells;

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(d) comparing the amount of binding of the substance to GPR54 in the test cells with the amount of binding of the substance to control cells that have not been transfected with GPR54;

wherein if the amount of binding of the substance is greater in the test cells as compared to the control cells, the substance is capable of binding to GPR54. Determining whether the substance is an agonist or antagonist can then be accomplished by the use of functional assays such as, e.g., the assay involving the use of promiscuous G-proteins described below.

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The conditions under which step (b) of the method is practiced are conditions that are typically used in the art for the study of protein-ligand interactions: e.g., physiological pH; salt conditions such as those represented by such commonly used buffers as PBS or in tissue culture media; a temperature of about 4°C to about 55°C.

In a particular embodiment of the above-described methods, the cells are eukaryotic cells. In another embodiment, the cells are mammalian cells. In other 15 embodiments, the cells are L cells L-M(TK-) (ATCC CCL 1.3), L cells L-M (ATCC CCL 1.2), HEK 293 (ATCC CRL 1573), Raji (ATCC CCL 86), CV-1 (ATCC CCL 70), COS-1 (ATCC CRL 1650), COS-7 (ATCC CRL 1651), CHO-K1 (ATCC CCL 61), 3T3 (ATCC CCL 92), NIH/3T3 (ATCC CRL 1658), HeLa (ATCC CCL 2), C127I (ATCC CRL 1616), BS-C-1 (ATCC CCL 26) or MRC-5 (ATCC CCL 171).

The assays described above can be carried out with cells that have been transiently or stably transfected with GPR54. Transfection is meant to include any method known in the art for introducing GPR54 into the test cells. For example, transfection includes calcium phosphate or calcium chloride mediated transfection. lipofection, infection with a retroviral construct containing GPR54, and electroporation.

Where binding of the substance or ligand to GPR54 is measured, such binding can be measured by employing a labeled substance or ligand. The substance or ligand can be labeled in any convenient manner known to the art, e.g., radioactively, fluorescently, enzymatically.

In particular embodiments of the above-described methods, GPR54 has an amino acid sequence of SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

The above-described methods can be modified in that, rather than exposing the test cells to the substance, membranes can be prepared from the test cells and those membranes can be exposed to the substance. Such a modification utilizing

membranes rather than cells is well known in the art and is described in, e.g., Hess et al., 1992, Biochem. Biophys. Res. Comm. 184:260-268.

Accordingly, the present invention provides a method for determining whether a substance is capable of binding to GPR54 comprising:

(a) providing test cells by transfecting cells with an expression vector that directs the expression of GPR54 in the cells;

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- (b) preparing membranes containing GPR54 from the test cells and exposing the membranes to a ligand of GPR54 under conditions such that the ligand binds to the GPR54 in the membranes;
- (c) subsequently or concurrently to step (b), exposing the membranes from the test cells to a substance;
- (d) measuring the amount of binding of the ligand to the GPR54 in the membranes in the presence and the absence of the substance;
- (e) comparing the amount of binding of the ligand to GPR54 in the membranes in the presence and the absence of the substance where a decrease in the amount of binding of the ligand to GPR54 in the membranes in the presence of the substance indicates that the substance is capable of binding to GPR54.

In particular embodiments, GPR54 has an amino acid sequence of SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

The present invention provides a method for determining whether a substance is capable of binding to GPR54 comprising:

- (a) providing test cells by transfecting cells with an expression vector that directs the expression of GPR54 in the cells;
- (b) preparing membranes containing GPR54 from the test cells and exposing the membranes from the test cells to the substance;
- (c) measuring the amount of binding of the substance to the GPR54 in the membranes from the test cells;
- (d) comparing the amount of binding of the substance to GPR54 in the membranes from the test cells with the amount of binding of the substance to membranes from control cells that have not been transfected with GPR54;

where if the amount of binding of the substance to GPR54 in the membranes from the test cells is greater than the amount of binding of the substance to the membranes from the control cells, then the substance is capable of binding to GPR54.

In particular embodiments, GPR54 has an amino acid sequence of SEO.ID.NO.:3 or SEQ.ID.NO.:4.

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As a further modification of the above-described methods, RNA encoding GPR54 can be prepared, e.g., by in vitro transcription using a plasmid containing GPR54 under the control of a bacteriophage T7 promoter, and the RNA can be microinjected into Xenopus oocytes in order to cause the expression of GPR54 in the oocytes. Substances are then tested for binding to the GPR54 expressed in the oocytes. Alternatively, rather than detecting binding, the effect of the substances on the electrophysiological properties of the oocytes can be determined.

The present invention includes assays by which GPR54 agonists and antagonists may be identified by their ability to stimulate or antagonize a functional response mediated by GPR54. One skilled in the art would be familiar with a variety of methods of measuring the functional responses of G-protein coupled receptors (see, e.g., Lerner, 1994, Trends Neurosci. 17:142-146 [changes in pigment distribution in melanophore cells]; Yokomizo et al., 1997, Nature 387:620-624 [changes in cAMP or calcium concentration; chemotaxis]; Howard et al., 1996, Science 273:974-977 [changes in membrane currents in *Xenopus* oocytes]; McKee et al., 1997, Mol. Endocrinol. 11:415-423 [changes in calcium concentration measured using the aequorin assay]; Offermanns & Simon, 1995, J. Biol. Chem. 270:15175, 15180 [changes in inositol phosphate levels]).

Accordingly, the present invention provides a method of identifying agonists and antagonists of GPR54 comprising:

- (a) providing test cells by transfecting cells with an expression vector that directs the expression of GPR54 in the cells;
- (b) exposing the test cells to a substance that is suspected of being an agonist or an antagonist of GPR54;
- (c) measuring the amount of a functional response of the test cells that have been exposed to the substance;
- (d) comparing the amount of the functional response exhibited by
 the test cells with the amount of the functional response exhibited by control cells;
 wherein if the amount of the functional response exhibited by the test
 cells differs from the amount of the functional response exhibited by the control cells,
 the substance is an agonist or antagonist of GPR54;

where the control cells are cells that have not been transfected with GPR54 but have been exposed to the substance or are test cells that have not been exposed to the substance.

In particular embodiments, GPR54 has an amino acid sequence of SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

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In particular embodiments, the functional response is selected from the group consisting of: changes in pigment distribution in melanophore cells; changes in cAMP or calcium concentration; changes in membrane currents in *Xenopus* oocytes; and changes in inositol phosphate levels.

GPR54 belongs to the class of proteins known as G-protein coupled receptors (GPCRs). GPCRs transmit signals across cell membranes upon the binding of ligand. The ligand-bound GPCR interacts with a heterotrimeric G-protein, causing the $G\alpha$ subunit of the G-protein to disassociate from the $G\beta$ and $G\gamma$ subunits. The $G\alpha$ subunit can then go on to activate a variety of second messenger systems.

Generally, a particular GPCR is only coupled to a particular type of G-protein. Thus, to observe a functional response from the GPCR, it is necessary to ensure that the proper G-protein is present in the system containing the GPCR. It has been found, however, that there are certain G-proteins that are "promiscuous." These promiscuous G-proteins will couple to, and thus transduce a functional signal from, virtually any GPCR. See Offermanns & Simon, 1995, J. Biol. Chem. 270:15175, 15180 (Offermanns). Offermanns described a system in which cells are transfected with expression vectors that result in the expression of one of a large number of GPCRs as well as the expression of one of the promiscuous G-proteins $G\alpha15$ or $G\alpha16$. Upon the addition of an agonist of the GPCR to the transfected cells, the GPCR was activated and was able, via $G\alpha15$ or $G\alpha16$, to activate the β isoform of phospholipase C, leading to an increase in inositol phosphate levels in the cells.

Therefore, by making use of these promiscuous G-proteins as in Offermanns, it is possible to set up functional assays for GPR54, even in the absence of knowledge of the G-protein with which GPR54 is coupled *in vivo*. One possibility is to create a fusion or chimeric protein composed of the extracellular and membrane spanning portion of GPR54 fused to a promiscuous G-protein. Such a fusion protein would be expected to transduce a signal following binding of ligand to the GPR54 portion of the fusion protein. Accordingly, the present invention provides a method of identifying antagonists of GPR54 comprising:

(a) providing cells that expresses a chimeric GPR54 protein fused at its C-terminus to a promiscuous G-protein;

- (b) exposing the cells to an agonist of GPR54;
- (c) subsequently or concurrently to step (b), exposing the cells to a substance that is a suspected antagonist of GPR54;
 - (d) measuring the level of inositol phosphates in the cells; where a decrease in the level of inositol phosphates in the cells in the presence of the substance as compared to the level of inositol phosphates in the cells in the absence of the substance indicates that the substance is an antagonist of GPR54.

Another possibility for utilizing promiscuous G-proteins in connection with GPR54 includes a method of identifying agonists of GPR54 comprising:

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GPR54;

- (a) providing cells that expresses both GPR54 and a promiscuous G-protein;
 - (b) exposing the cells to a substance that is a suspected agonist of
- (c) measuring the level of inositol phosphates in the cells; where an increase in the level of inositol phosphates in the cells as compared to the level of inositol phosphates in the cells in the absence of the suspected agonist indicates that the substance is an agonist of GPR54.

Levels of inositol phosphates can be measured by monitoring calcium mobilization. Intracellular calcium mobilization is typically assayed in whole cells under a microscope using fluorescent dyes or in cell suspensions via luminescence using the aequorin assay.

In a particular embodiment of the above-described method, the cells are eukaryotic cells. In another embodiment, the cells are mammalian cells. In other embodiments, the cells are L cells L-M(TK⁻) (ATCC CCL 1.3), L cells L-M (ATCC CCL 1.2), 293 (ATCC CRL 1573), Raji (ATCC CCL 86), CV-1 (ATCC CCL 70), COS-1 (ATCC CRL 1650), COS-7 (ATCC CRL 1651), CHO-K1 (ATCC CCL 61), 3T3 (ATCC CCL 92), NIH/3T3 (ATCC CRL 1658), HeLa (ATCC CCL 2), C1271 (ATCC CRL 1616), BS-C-1 (ATCC CCL 26), MRC-5 (ATCC CCL 171), Xenopus oocytes, or Xenopus melanophores.

In a particular embodiment of the above-described method, the cells are transfected with expression vectors that direct the expression of GPR54 and the promiscuous G-protein in the cells.

The conditions under which step (b) of the method is practiced are conditions that are typically used in the art for the study of protein-ligand interactions: e.g., physiological pH; salt conditions such as those represented by such commonly used buffers as PBS or in tissue culture media; a temperature of about 4°C to about 55°C.

In a particular embodiment of the above-described method, the promiscuous G-protein is selected from the group consisting of $G\alpha15$ or $G\alpha16$. Expression vectors containing $G\alpha15$ or $G\alpha16$ are known in the art. See, e.g., Offermanns; Buhl et al., 1993, FEBS Lett. 323:132-134; Amatruda et al., 1993, J. Biol. Chem. 268:10139-10144.

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The above-described assay can be modified to form a method to identify antagonists of GPR54. Such a method is also part of the present invention and comprises:

- (a) providing cells that expresses both GPR54 and a promiscuous 15 G-protein;
 - (b) exposing the cells to a substance that is an agonist of GPR54;
 - (c) subsequently or concurrently to step (b), exposing the cells to a substance that is a suspected antagonist of GPR54;
 - (d) measuring the level of inositol phosphates in the cells; where a decrease in the level of inositol phosphates in the cells in the presence of the suspected antagonist as compared to the level of inositol phosphates in the cells in the absence of the suspected antagonist indicates that the substance is an antagonist of GPR54.

In a particular embodiment of the above-described method, the cells
are eukaryotic cells. In another embodiment, the cells are mammalian cells. In other
embodiments, the cells are L cells L-M(TK⁻) (ATCC CCL 1.3), L cells L-M (ATCC
CCL 1.2), 293 (ATCC CRL 1573), Raji (ATCC CCL 86), CV-1 (ATCC CCL 70),
COS-1 (ATCC CRL 1650), COS-7 (ATCC CRL 1651), CHO-K1 (ATCC CCL 61),
3T3 (ATCC CCL 92), NIH/3T3 (ATCC CRL 1658), HeLa (ATCC CCL 2), C1271

(ATCC CRL 1616), BS-C-1 (ATCC CCL 26), MRC-5 (ATCC CCL 171), Xenopus
oocytes, or Xenopus melanophores.

The conditions under which steps (b) and (c) of the method are practiced are conditions that are typically used in the art for the study of protein-ligand interactions: e.g., physiological pH; salt conditions such as those represented by such

commonly used buffers as PBS or in tissue culture media; a temperature of about 4°C to about 55°C.

In a particular embodiment of the above-described method, the cells are transfected with expression vectors that direct the expression of GPR54 and the promiscuous G-protein in the cells.

In a particular embodiment of the above-described method, the promiscuous G-protein is selected from the group consisting of Ga15 or Ga16.

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In particular embodiments of the above-described methods, GPR54 has an amino acid sequence of SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

While the above-described methods are explicitly directed to testing whether "a" substance is an agonist or antagonist of GPR54, it will be clear to one skilled in the art that such methods can be adapted to test collections of substances, e.g., combinatorial libraries, to determine whether any members of such collections are activators or inhibitors of GPR54. Accordingly, the use of collections of substances, or individual members of such collections, as the substance in the above-described methods is within the scope of the present invention.

Agonists and antagonists of GPR54 that are identified by the above-described methods are expected to have utility in the treatment of diseases that involve the inappropriate expression of GPR54. In particular, given the resemblance between GPR54 and the galanin receptors, it is expected that agonists and antagonists of GPR54 will have pharmacological activity and be useful in a manner similar to that in which agonists and antagonists of the galanin receptors are useful. Therefore, agonists and antagonists of GPR54 are expected to be useful in the treatment of: eating disorders and obesity; Alzheimer's disease and other disorders affecting memory; pain; sexual disorders; and growth hormone imbalances.

The present invention includes pharmaceutical compositions comprising agonists and antagonists of GPR54. The agonists and antagonists are generally combined with pharmaceutically acceptable carriers to form pharmaceutical compositions. Examples of such carriers and methods of formulation of pharmaceutical compositions containing agonists and antagonists and carriers can be found in Remington's Pharmaceutical Sciences. To form a pharmaceutically acceptable composition suitable for effective administration, such compositions will contain a therapeutically effective amount of the agonists and antagonists.

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Therapeutic or prophylactic compositions are administered to an individual in amounts sufficient to treat or prevent conditions where GPR54 activity is abnormal. The effective amount can vary according to a variety of factors such as the individual's condition, weight, gender, and age. Other factors include the mode of administration. The appropriate amount can be determined by a skilled physician.

Compositions can be used alone at appropriate dosages. Alternatively, co-administration or sequential administration of other agents can be desirable.

The compositions can be administered in a wide variety of therapeutic dosage forms in conventional vehicles for administration. For example, the compositions can be administered in such oral dosage forms as tablets, capsules (each including timed release and sustained release formulations), pills, powders, granules, elixirs, tinctures, solutions, suspensions, syrups and emulsions, or by injection. Likewise, they can also be administered in intravenous (both bolus and infusion), intraperitoneal, subcutaneous, topical with or without occlusion, or intramuscular form, all using forms well known to those of ordinary skill in the pharmaceutical arts.

Advantageously, compositions can be administered in a single daily dose, or the total daily dosage can be administered in divided doses of two, three or four times daily. Furthermore, compositions can be administered in intranasal form via topical use of suitable intranasal vehicles, or via transdermal routes, using those forms of transdermal skin patches well known to those of ordinary skill in that art. To be administered in the form of a transdermal delivery system, the dosage administration will, of course, be continuous rather than intermittent throughout the dosage regimen.

The dosage regimen utilizing the compositions is selected in accordance with a variety of factors including type, species, age, weight, sex and medical condition of the patient; the severity of the condition to be treated; the route of administration; the renal, hepatic and cardiovascular function of the patient; and the particular composition thereof employed. A physician of ordinary skill can readily determine and prescribe the effective amount of the composition required to prevent, counter or arrest the progress of the condition. Optimal precision in achieving concentrations of composition within the range that yields efficacy without toxicity requires a regimen based on the kinetics of the composition's availability to target sites. This involves a consideration of the distribution, equilibrium, and elimination of a composition.

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The present invention also includes methods of expressing GPR54 in recombinant systems and then utilizing the recombinantly expressed GPR54 receptor protein for counter-screening. When screening compounds in order to identify potential pharmaceuticals that specifically interact with a target receptor, it is necessary to ensure that the compounds identified are as specific as possible for the target receptor. To do this, it is necessary to screen the compounds against as wide an array as possible of receptors that are similar to the target receptor. Thus, in order to find compounds that are potential pharmaceuticals that interact with receptor A, it is necessary not only to ensure that the compounds interact with receptor A (the "plus target") and produce the desired pharmacological effect through receptor A, it is also necessary to determine that the compounds do not interact with receptors B, C, D, etc. (the "minus targets"). In general, as part of a screening program, it is important to have as many minus targets as possible (see Hodgson, 1992, Bio/Technology 10:973-980, at 980). Therefore, GPR54 proteins and DNA encoding GPR54 proteins have utility in counter-screens. That is, they can be used as "minus targets" in counterscreens in connection with screening projects designed to identify compounds that specifically interact with other G-protein coupled receptors.

The DNA of the present invention, or hybridization probes based upon the DNA, can be used in chromosomal mapping studies in order to identify the precise chromosomal location of the GPR54 gene or of genes encoding proteins related to GPR54. While the present inventors have determined that the human GPR54 gene is located at chromosome 19p13.3, it may be desirable to perform mapping studies to even more precisely locate the human GPR54 gene. Such mapping studies can be carried out using well-known genetic and/or chromosomal mapping techniques such as, e.g., linkage analysis with respect to known chromosomal markers or in situ hybridization. See, e.g., Verma et al., 1988, Human Chromosomes: A Manual of Basic Techniques, Pergamon Press, New York, NY. After identifying the precise chromosomal location of the GPR54 gene or genes encoding proteins related to GPR54, this information can be compared with the locations of known diseasecausing genes contained in genetic map data (such as the data found in the genome issue of Science (1994, 265:1981-2144). In this way, one can correlate the chromosomal location of the GPR54 gene or of genes encoding proteins related to GPR54 with the locations of known disease-causing genes and thus help to limit the region of DNA containing such disease-causing genes. This will simplify the process of cloning such disease-causing genes. Also, once linkage between the precise

chromosomal location of the GPR54 gene or of genes encoding proteins related to GPR54 and the locations of a known disease-causing gene is established, that linkage can be used diagnostically to identify restriction fragment length polymorphisms (RFLPs) in the vicinity of the GPR54 gene or of genes encoding proteins related to GPR54. Such RFLPs will be associated with the disease-causing gene and thus can be used to identify individuals carrying the disease-causing gene.

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For such chromosomal mapping studies as described herein, it may be advantageous to use, in addition to the DNA of the present invention, the reverse complement of the DNA of the present invention or RNA corresponding to the DNA of the present invention.

Nucleotide sequences that are complementary to the GPR54 sequences disclosed herein can be synthesized for use in antisense therapy. Such antisense molecules can be DNA, stable derivatives of DNA such as phosphorothioates or methyl phosphonates, RNA, stable derivatives of RNA such as 2'-O-alkyl RNA, or other forms of GPR54 antisense molecules. GPR54 antisense molecules can be introduced into cells by a variety of methods, e.g., microinjection, liposome encapsulation, or by expression from vectors harboring the antisense sequence. GPR54 antisense therapy is expected to be particularly useful in the treatment of conditions where it is beneficial to reduce GPR54 activity.

The present invention also includes antibodies to the GPR54 protein. Such antibodies may be polyclonal antibodies or monoclonal antibodies and are useful in treating disorders that involve the inappropriate expression or activity of the GPR54 protein. The antibodies of the present invention are raised against the entire GPR54 protein or against suitable antigenic fragments of the protein that are coupled to suitable carriers, e.g., serum albumin or keyhole limpet hemocyanin, by methods well known in the art. Methods of identifying suitable antigenic fragments of a protein are known in the art. See, e.g., Hopp & Woods, 1981, Proc. Natl. Acad. Sci. USA 78:3824-3828; and Jameson & Wolf, 1988, CABIOS (Computer Applications in the Biosciences) 4:181-186.

For the production of polyclonal antibodies, GPR54 protein or an antigenic fragment, coupled to a suitable carrier, is injected on a periodic basis into an appropriate non-human host animal such as, e.g., rabbits, sheep, goats, rats, mice. The animals are bled periodically and sera obtained are tested for the presence of antibodies to the injected antigen. The injections can be intramuscular, intraperitoneal, subcutaneous, and the like, and can be accompanied with adjuvant.

For the production of monoclonal antibodies, GPR54 protein or an antigenic fragment, coupled to a suitable carrier, is injected into an appropriate non-human host animal as above for the production of polyclonal antibodies. In the case of monoclonal antibodies, the animal is generally a mouse. The animal's spleen cells are then immortalized, often by fusion with a myeloma cell, as described in Kohler & Milstein, 1975, Nature 256:495-497. For a fuller description of the production of monoclonal antibodies, see Antibodies: A Laboratory Manual, Harlow & Lane, eds., Cold Spring Harbor Laboratory Press, 1988.

Gene therapy may be used to introduce GPR54 polypeptides into the cells of target organs. Nucleotides encoding GPR54 polypeptides can be ligated into viral vectors which mediate transfer of the nucleotides by infection of recipient cells. Suitable viral vectors include retrovirus, adenovirus, adeno-associated virus, herpes virus, vaccinia virus, and polio virus based vectors. Alternatively, nucleotides encoding GPR54 polypeptides can be transferred into cells for gene therapy by non-viral techniques including receptor-mediated targeted transfer using ligand-nucleotide conjugates, lipofection, membrane fusion, or direct microinjection. These procedures and variations thereof are suitable for *ex vivo* as well as *in vivo* gene therapy. Gene therapy with GPR54 polypeptides will be particularly useful for the treatment of diseases where it is beneficial to elevate GPR54 activity.

A cDNA fragment encoding full-length GPR54 can be isolated from an appropriate human cDNA library by using the polymerase chain reaction (PCR) employing suitable primer pairs. Such primer pairs can be selected based upon the cDNA sequence for GPR54 shown in Figure 1 as SEQ.ID.NO.:1. Suitable primer pairs would be, e.g.:

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5'-ATG CAC ACC GTG GCT ACG TCC-3' (SEQ.ID.NO.:11) and 5'-TCA GAG AGG GGC GTT GTC CTC-3' (SEQ.ID.NO.:12).

The above primers may contain restriction sites in their 5' ends to facilitate cloning of the amplified cDNA into suitable vectors, e.g., pcDNA3.1. The above primers are meant to be illustrative. One skilled in the art would recognize that a variety of other suitable primers can be designed.

PCR reactions can be carried out with a variety of thermostable enzymes including but not limited to AmpliTaq, AmpliTaq Gold, or Vent polymerase. For AmpliTaq, reactions can be carried out in 10 mM Tris-Cl, pH 8.3, 2.0 mM

MgCl₂, 200 μM for each dNTP, 50 mM KCl, 0.2 μM for each primer, 10 ng of DNA template, 0.05 units/μl of AmpliTaq. The reactions are heated at 95°C for 3 minutes and then cycled 35 times using the cycling parameters of 95°C, 20 seconds, 62°C, 20 seconds, 72°C, 3 minutes. In addition to these conditions, a variety of suitable PCR protocols can be found in PCR Primer, A Laboratory Manual, edited by C.W. Dieffenbach and G.S. Dveksler, 1995, Cold Spring Harbor Laboratory Press; or PCR Protocols: A Guide to Methods and Applications, Michael et al., eds., 1990, Academic Press.

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A suitable cDNA library from which a clone encoding GPR54 can be isolated would be a human cDNA library made from RNA from brain tissue. Such libraries can be prepared by methods well-known in the art. Alternatively, several commercially available libraries would be suitable, e.g., cDNA libraries such as human fetal brain, catalog #937227 from Stratagene, Inc., La Jolla, CA, USA, and human brain hypothalamus, catalog #HL1172a, from Clontech Laboratories, Inc., Palo Alto, CA, USA. The primary clones of such libraries can be subdivided into pools with each pool containing approximately 20,000 clones and each pool can be amplified separately.

By this method, a cDNA fragment encoding an open reading frame of 398 amino acids (SEQ.ID.NO.:3) can be obtained. This cDNA fragment can be cloned into a suitable cloning vector or expression vector. For example, the fragment can be cloned into the mammalian expression vector pcDNA3.1 (Invitrogen, San Diego, Ca). GPR54 protein can then be produced by transferring an expression vector encoding GPR54 into suitable host cells and growing the host cells under appropriate conditions. GPR54 protein can then be isolated by methods well known in the art.

As an alternative to the above-described PCR method, a cDNA clone encoding GPR54 can be isolated from a cDNA library using as a probe oligonucleotides specific for GPR54 and methods well known in the art for screening cDNA libraries with oligonucleotide probes. Such methods are described in, e.g., Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual; Cold Spring Harbor Laboratory, Cold Spring Harbor, New York; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K., Vol. I, II. Oligonucleotides that are specific for GPR54 and that can be used to screen cDNA libraries can be readily designed based upon the cDNA sequence of GPR54 shown in Figure 1 as SEO.ID.NO.:1 and can be synthesized by methods well-known in the art.

Genomic clones containing the GPR54 gene can be obtained from commercially available human PAC or BAC libraries, e.g., from Research Genetics, Huntsville, AL. Alternatively, one may prepare genomic libraries, for example in P1 artificial chromosome vectors, from which genomic clones containing the GPR54 can be isolated, using probes based upon the GPR54 nucleotide sequences disclosed herein. Methods of preparing such libraries are known in the art (Ioannou et al., 1994, Nature Genet. 6:84-89).

The following non-limiting examples are presented to better illustrate the invention.

EXAMPLE 1

PCR amplification and cDNA library screening

A rat brain 5' Stretch cDNA library (Clontech) was amplified by the polymerase chain reaction (PCR) using proof-reading *Pfu* polymerase (Stratagene) and degenerate oligonucleotides based upon sequences encoding GPCR conserved transmembrane (TM) region 3

P1: 5'-CTGACCGGCATGABDETFGADCGHTA-3' (SEQ.ID.NO.:9)

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and transmembrane (TM) region 7

P2: 5'-GAAGGCGTAGAFBAIJGGKTT) -3' (SEQ.ID.NO.:10)

where B = C or G, D = C or G, E = A or G or G, E = C or G or G, E = A or G.

PCR conditions were as follows: denaturation at 94°C for 30 sec, annealing at 55, 48, 45, 42, or 40°C for 40 sec, and extension at 72°C for 30 sec, for 30 cycles, followed by a 7 min extension at 72°C. The PCR products were extracted with phenol/chloroform, precipitated with ethanol and electrophoresed on a low melting point agarose gel. PCR product bands in the expected size range were excised from the gel, ligated into the *EcoRV* site of pBluescript SK(-) (Stratagene)

and sequenced. One insert appeared to encode a novel GPCR and was labeled with [32P] dCTP-α (NEN) by nick translation (Amersham) and used to screen the same library amplified above as previously described (Marchese et al., 1994, Genomics 23:609-618). Positive phage clones were plaque purified and their inserts amplified by PCR using *Pfu* polymerase and primers flanking the lgt11 *EcoR*I cloning site. The PCR products were blunt-end ligated into the *EcoRV* site of pBluescript SK(-) (Stratagene) and sequenced on both strands.

EXAMPLE 2

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Rat mRNAs from several rat tissues were extracted as described previously (Marchese et al., 1994, Genomics 23:609-618). Briefly, total RNA was extracted by the method of Chomczynski & Sacchi, 1987, Anal. Biochem. 162:156-159 and poly (A)⁺ RNA isolated using oligo(dT) cellulose spin columns (Pharmacia, Uppsala, Sweden). RNA was denatured and size fractionated on a 1% formaldehyde agarose gel, transferred onto nylon membrane and immobilized by UV irradiation. The blots were hybridized with a 32P-labeled DNA fragment encoding GPR54, washed with 2X SSPE and 0.1% SDS at 50°C for 20 min and again with 0.1X SSPE and 0.1% SDS at 50°C for 2 h and exposed to X-ray film at -70°C in the presence of an intensifying screen.

EXAMPLE 3

In situ hybridization analysis

An 35S-labeled DNA fragment encoding GPR54 was used as a probe for *in situ* hybridization. Preparation of rat brain sections and *in situ* hybridization procedures were done as previously described (O'Dowd et al., 1996, FEBS Lett 394:325-329).

EXAMPLE 4

Expression of GPR54 cDNA in COS-7 mammalian cells

The African Green Monkey SV40 transformed kidney cell line (COS-7 cells), obtained from the American Type Culture Collection, was grown in

5 Dulbecco's modified Eagle's medium supplemented with 10% heat-inactivated fetal calf serum (Sigma), 50 units/ml penicillin, 50 μg/ml streptomycin (Flow Laboratories, McLean, VA), and 2 mM glutamine (Flow Laboratories) at 37°C under an atmosphere of 6% CO2. 5 X 106 cells per 175- cm² culture flask were seeded in 20 ml of media and transiently transfected at 80% confluence with either 2.75, 5.5, or 11.65 μg of pcDNA3-GPR54 or pcIneo-hGALR1 plasmids and 70 μl of LipofectAMINE reagent (Life Technologies, Inc.), following recommendations of the manufacturer. Two days after transfection, cells were harvested following dissociation in enzyme-free dissociation solution (Specialty Media, Lavallette, NJ).

EXAMPLE 5

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Membrane preparation and radioligand binding assays

Membranes were prepared from transfected cells by disruption by pressurized nitrogen cavitation in ice-cold membrane buffer (10 mM Tris, pH 7.4, 10 mM phenylmethylsulfonylfluoride, 10 mM phosphoramidon). After a low speed (1100 x g for 10 min. at 4°C) and a high speed centrifugation (38,700 x g for 15 min. 20 at 4°C), membranes were resuspended in buffer and their protein concentration determined (Bio-Rad assay kit). Binding of 125I-human galanin (specific activity of 2200 Ci/mmol, DuPont NEN) was measured in membranes using a buffer of 25 mM Tris, pH 7.4, 0.3% BSA, 2 mM MgCl₂, 4 mg/ml phosphoramidon, and 10 mM leupeptin in a total volume of 250 ml. 200 pM of ¹²⁵I-human galanin was used. 25 Reactions were initiated by the addition of membranes and the incubation was allowed to proceed at room temperature for 2 hours. Non-specific binding was defined as the amount of radioactivity remaining bound in the presence of 10 mM unlabeled human galanin. Incubations were terminated by rapid filtration through GF/C filters which had been presoaked with 0.1% polyethylamine using a TOMTEC 30 (Orange, CT) cell harvester.

The present invention is not to be limited in scope by the specific embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description. Such modifications are intended to fall within the scope of the appended claims.

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Various publications are cited herein, the disclosures of which are incorporated by reference in their entireties.

WHAT IS CLAIMED:

1. An isolated DNA comprising nucleotides encoding a polypeptide having the amino acid sequence SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

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- 2. The DNA molecule of claim 1 comprising a nucleotide sequence selected from the group consisting of SEQ.ID.NO.:1; positions 1-1,194 of SEQ.ID.NO.:1; SEQ.ID.NO.:2; and positions 61-1,245 of SEQ.ID.NO.:2.
- 10 3. A DNA molecule that hybridizes under stringent conditions to the DNA of claim 1.
 - 4. An expression vector comprising the DNA of claim 1.

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- 5. A recombinant host cell comprising the DNA of claim 1.
- 6. An isolated polypeptide comprising a GPR54 protein having the amino acid sequence SEQ.ID.NO.:3 or SEQ.ID.NO.:4

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- 7. The isolated polypeptide of claim 6 that is substantially free from other proteins.
- 8. The isolated polypeptide of claim 6 containing a single amino acid substitution.
 - 9. The isolated polypeptide of claim 6 containing two or more amino acid substitutions where the substitutions are conservative and do not occur in positions where GPR54 and any of the rat GALR1, GALR2, or GALR3 receptors share the same amino acid.
 - 10. A method for determining whether a substance is an agonist or antagonist of GPR54 comprising:
 - (a) transfecting cells with an expression vector encoding GPR54;

(b) allowing the transfected cells to grow for a time sufficient to allow GPR54 to be expressed;

- (c) exposing the cells to a labeled ligand of GPR54 in the presence and in the absence of the substance;
- (d) measuring the binding of the labeled ligand to GPR54; where if the amount of binding of the labeled ligand is less in the presence of the substance than in the absence of the substance, then the substance is an agonist or antagonist of GPR54;

where GPR54 has the amino acid sequence SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

- 11. A method for determining whether a substance is capable of binding to GPR54 comprising:
- (a) providing test cells by transfecting cells with an expression vector that directs the expression of GPR54 in the cells;
 - (b) exposing the test cells to the substance;
 - (c) measuring the amount of binding of the substance to GPR54 in the test cells;
- (d) comparing the amount of binding of the substance to GPR54 in the test cells with the amount of binding of the substance to control cells that have not been transfected with GPR54;

wherein if the amount of binding of the substance is greater in the test cells as compared to the control cells, the substance is capable of binding to GPR54; where GPR54 has the amino acid sequence SEQ.ID.NO.:3 or

25 SEQ.ID.NO.:4.

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- 12. A method of identifying agonists and antagonists of GPR54 comprising:
- (a) providing test cells by transfecting cells with an expression vector that directs the expression of GPR54 in the cells;
 - (b) exposing the test cells to a substance that is suspected of being an agonist or an antagonist of GPR54;
 - (c) measuring the amount of a functional response of the test cells that have been exposed to the substance;

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(d) comparing the amount of the functional response exhibited by the test cells with the amount of the functional response exhibited by control cells; wherein if the amount of the functional response exhibited by the test cells differs from the amount of the functional response exhibited by the control cells, the substance is an agonist or antagonist of GPR54;

where the control cells are cells that have not been transfected with GPR54 but have been exposed to the substance or are test cells that have not been exposed to the substance;

where GPR54 has the amino acid sequence SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

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13. An antibody that binds specifically to GPR54 where GPR54 has the amino acid sequence SEQ.ID.NO.:3 or SEQ.ID.NO.:4.

A CC	Val GTG 120	Ser 166 183	Ala GCC 246	Tyr TAC 309	Val GTC 372
Asn AAC 54	Ald	Asn	I i e ATC	Leu	GIn CAG
A CCC	Arg CGG	G1y 666	Tyr	Leu CTG	GIn
5 5 8	Pro 111	Val GTG 174	Phe TTC 237	Ald GCC 300	11e ATC 363
Ala GCA 45	Ser TCG	Leu CTG	Asn	Thr	Tyr TAC
6 <u>1</u> 9	Pro CCT	65. 66.	Thr	Phe TTC	Asn
7rp 166		Leu CTG 165	Va GTG 228	Pro CCC 291	Val GTC 354
Ser 36	Pro	Leu CTG	Thr	Val GTC	Phe TTC
P G G G	61y 660	Met ATG	Arg CCC	Cys 760	Lys AAG
Asn	Asp GAC 93	Leu CTG 156	Met ATG 219	Cys TGC 282	Cys TGC 345
Pro CCC 27	Ser TCC	A SS	£8	Leu CTG	Met ATG
GIY GGA	Ald GCC	Ald SS SS	Lys AAG	Leu	Phe TTC
Ser	Asn AAC 84	Phe TTC 147	His CAC 210	Phe TTC 273	Asp GAC 336
Thr ACG 18	Ald	Phe TTC	Arg CGC	Thr ACC	61.y 900
Ala GCT	61y 660	Leu	Cys TGC	Val GTG	Leu CTG
Val GTG	Cys TGT 75	Pro CCC 138	11e ATC 201	Asp GAC 264	Val GTG 327
ACC 9	61y 660	Val GTG	Val GTC	Thr ACG	175 135
His CAC	Pr SS	Leu	Tyr TAC	Ala	<u>၄၂</u> ၁ ၁၁၁
Met ATG	Cys TGC 66	Trp TGC 129	11e ATC 192	Ala GGG 225	Pro CCC 318
	61y 660	Ala	Val	Leu CTG	Leu CTG
	Ser TCC	Asp GAC	Leu CTG	Asn	5 5 5 5 5 5 5

FIG. 1A

Val GTG 435	11e ATC 498	Pro CCC 561	Asn AAC 624	Arg CGC 687	Leu CTG 750	Leu CTC 813
	Ser AGC	G1y GGG	Tyr TAC	Leu CTG	Val GTG	Leu CTG
Val GTG	Leu	Pro CCC	Leu CTG	Met ATG	GIN	Val
Tyr TAC 426	Ser AGC 489	Ser TCA 552	Ala GCA 615	Ala GCC 678	G1y GGG 741	
	Val GTC	Leu CTG	Phe TTC	Ala GCG	Gin	A GCC
	Ala	Arg CCC	A CCC	Tyr TAT	Leu CTG	A GCG
Asp GAC 417		His CAC 543	Arg 060 606	Cys TGC 669	A10 GCC 732	Va GTG 795
Val GTG	Ald GCG	Leu CTG	G l u GAG	A CCC	Ser AGC	
Ser	Leu	Ala	Leu	Cys 160	Asp GAT	Arg 066
Met ATG 408	Arg CCC 471	Leu CTC 534	Ala GCC 597	Thr ACC 660		
A d CCC	Pro CCC	Val GTG	Arg CGC	A La GCC	5 5 5 5	
ACC	Thr ACG	58	Ser AGC	Leu CTC	A GCC	Lys AAG
Leu CTG 399	Arg CCC 462	A1g 606 525	Pro CCC 588	Leu CTG 651	Pro CCC 714	Ala GCC 777
Thr	Arg CCC	Ser TCT	Phe TTC	P. 6	Arg OSC OSC	Arg CGG
Ala SCC	His	Val GTG	Ala GCC	Leu CTG	Val GTG	Val GTG
Cys TGT 390	Leu CTG 453	Ala 606 516	Glu GAG 579		Val Ala GTC GCC 705	A14 GCC 768
ACG ACG	Ala GCC	A la	Ser AGT	Tyr TAC	Val GTC	
A P GCC	Arg CCC	Ser TCT	Cys 160	Leu CTG	Arg 006	A GCA
Gln CAG 381	Leu TTG 444	G1y GGC 507	Tyr 1AC 570	Ala 600 633	G1y GGC 696	
Val GTG	5 5 5	Val GTA	9 A G	Leu CTG	Leu	GI u GAG
Ser TCG	Phe TTC	1rp 166	Arg CCC	Leu CTG	His	A GCA

61y 660 876	Tyr 1AC 939	Phe TTC 1002	Ser Asp TCG GAC 1065	His Ala Glu Leu Leu Arg Leu Gly Ser His Pro Ala Pro Ala Arg Ala Gln CAC GCG GAG CTG CTC CGC CTG GGG TCC CAC CCG GCC CCC GCC AGG GCG CAG 1083 1092 1101 1110	Pro 1191
Ala 600	Ser TCC	Ala	Ser TCG	A C	A GCC
5 5 5 5	Met Ser ATG TCC	GIn	P 55	Arg AGG	Asn AAC
G1y GGC 867	Cys 160 930	S CGA CAG GCC 993	G1y GGA 1056	A10 600 1119	Asp GAC 1182
Leu	His	Phe TTC	Pro Gly Pro ccc GGA CCC 1056	500°	CAG
Ala 606	Ala GCT	His	Arg CGG	A la	61 <u>y</u>
Gln CAG 858	Trp Ala His TGG GCT CAC 921	y Ser His Phe / C TCG CAC TTC (984	Arg CCC 1047	Pro 200 110	Leu CTG 173
Leu CTG	ACC	SC 200	5 5 1	His CAC	Val GTC
Val GTG	Lys Thr F AAG ACC	Leu CTG	Arg CCC	Ser TCC	Cys TGC
Phe Leu Val Leu Gln Ala Leu Gly Pro Ala TTC CTG GTG CTG CAG GCG CTG GGC CCC GCG 849 858 867	Leu CTT 912	Tyr Ala Phe	Pro Arg Arg Pro Arg Arg Pro Arg Arg CCG CCC CCC CCC CCC CCC CCC 1029 1038 1047	G19 G66 101	Leu CTG
Phe TTC	A C C C C C C C C C C C C C C C C C C C	Ala GCC	5 82_	Leu CTG	<u>≻</u> 88
Leu CTG	Arg Ser Tyr Ala Ala Tyr Ala CGC AGC TAC GCC GCC TAC GCG 894 903	Tyr TAC	Arg OGC	Arg CGC	Arg CGC
Trp Gly Pro Ile Gln Leu TGG GGC CCC ATC CAG CTG 831 840	Ala GCC 903	Leu CTC 966	Arg CGC 1029	Leu CTC 1092	A1a 666
I le ATC	Ala GCC	Leu CTG		Leu CTG	A CC
o C C C	Tyr	Pro	Ala 606	G I u	Leu CTG
61y 66C 831	Ser AGC 894	Asn AAC 957	Pro Cys Ala CCC TGC GCC 1020	His Alg CAC GCG 1083	66 66 46
Trp 766	Arg CGC	Leu Asn CTG AAC 957	org CC	His CAC	Ser AGT
Cys TGC	Pro CCA	A la	Cys 160	S C A	Ser AGC
Ala GCC 822	His CAC 885	Ser TCC 948	Arg Val CGC GTC 1011	Ala Ala GCA GCC 1074	Pro Gly CCA GGG
Ala	1rp 166	Asn	Arg GC 1	Ala GCA 1	SS.
Phe /	Ser TCC	Ser AGC	Arg CGC	S S S S	Lys AAG
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-16.1C

පු .	16C	CTG	CTT	GAT	CTG	ACT	၁၅၁	පු	CGT
AGG	TCC GCA 123		TTC 255	ACA 321	GTG 387		GC 519	600 585	AGC 651
GAC 54	32	၁၁၅	ATC	ည္တ	166	161	CAC	32	8
၁၁	GCT	GAT	GTC	33	ACC	ACA	CTT	GTT	Ë
QAG CAG	AAC 114	CTG 180	CTG 246	CTG 312	CCC 378	GCC 444	GCA 510	GCT 576	6CG 642
CCA CAG TCC CAG GAC GCA ATC TGT GAA GGC TGC CTG GAG GAG GAG GGC GAC AGG GCC GAC AGG GAC AGG GCC AGG GCC GAC AGG GCC AGG GCC GAC AGG GCC AGG GCC GAC AGG GCC AGG GCC AGG GCC GAC AGG GCC AGG AGG	GGT CCG AAC GTG AGC TGG TGG GCT CCG TCC AAC GCT 87 96 105 114	CCA AGG CCC CTG GAT GCC TGG 171 180 189	TCA	ACC GTC ACC AAT TTC TAC ATC GCT AAC CTG GCG GCC ACA 285 294 303 312 321	CTG	AAC TAC ATC CAG CAG GTC TCG GTG CAA GCC ACA TGT GCC 417 426 435 444 453	CTG CGT GCA CTT CAC 510	GGT TCC GCA GCT GTT TCC 567 576	CCT CAC ACC TAC TGC AGT GAG GCG TTT CCC 624 633 642
CAC CAC	8	AGG	AAC	CCT	99	CTC	CTG	32	AGT
O V C	GCT 105	17A	666 237	ATC 303	TAT 369	TCC 435	505	GGT 567	TGC 633
CTG 36	351	GAT GGC CCA GGC TCC GCG 153	GTC	TAC	CTC	GTC	110	AGC CTT AGC ATC TGG GTG 549 558	TAC
<u> </u>	16 6	32	CTA	110	CTC	CAG	GTG	351	λ C
ပ္ပ	AGC 96	GGC 162	GGG 228	AAT 294	360	CAG 426	ACT 492	ATC 558	CCT CAC ACC 624 FIC 2 A
GAA 27	GTG	SS	CTG	ACC	ACC	ATC	GTG	AGC	133
161	AAC	299	116	GTC	110	TAC	TAC	CTT	ccc 666 615
ATC	000 87	GAT 153	ATG 219	ACC 285	351	AAC 417	166 483	AGC 549	CCC 615
6CA 18	561	35	CTA	CAG	GTA	GTC	၁ဗ	GCC CTG ACT GTC 540	25
GAC	116	သွ	၁၁၅	ATG	760	110	GAC	ACT	CTG
CAG	ACG 78	AAT 144	GCT 210	CAC 276	TGC 342	AAA 408	GTG 474	CTG 540	ည 909
ည္ စ	ည္ဟ	GTC	110	AAG	CTG	760	AGT	ည္ဟ	CAC
CAG	GAG	GGT	III	CAC	CTT	ATG	ATG	CTG	СТС
CCA	SCA 69	TGC 135	CTG 201	000 267	TTC 333	TTC ATG TGC AAA TTC GTC 399 408	600 465) 0GC CTG 0 531	GCT CTG CAC CGC CTG TCG 597 606
	GCC GCA GAG GCG ACG TTG 69 78	GGC TGC GGT GTC AAT GCC TCC 135 144	CCC CTG TTT TTC GCT GCC CTA ATG TTG CTG GGG CTA GTC GGG AAC TCA CTG GTC ATC TTC 201 210 219 228 237 246 255	ATC TGC CGC CAC AAG CAC ATG CAG 267 276	GTC ACT TTC CTT CTG TGC TGC GTA CCC TTC ACC GCG CTC CTC TAT CCG CTG CCC TGC TGC 333 342 351 350 360 369 378	GGA GAC	TTG ACA GCC ATG AGT GTG GAC CGC TGG TAC GTG ACT GTG TTC 465 474 483 492	g	CTG
	ATG	83	616	ATC	CTC	GGA	116	ACT	GTG

CTGTGC ATG TCC TAC AGC AAT TCT GCG CTC AAC CCG CTG CTC TAT GCC TTC CTG GGT TCC CAC TTC AGA 1002 1002 1011 1020 1029 1039 1038 1047 CAG GCC TTC TGC CGC GTG TGC CCC TGC GGC CCG CAA CGC CAG CGT CGG CCC CAC GCG TCA GCG CAC 1059 1059 1104 1113 TOG GAC CGA GCC GCA CCC CAT AGT GTG COG CAC AGC CGG GCT GCG CAC CCT GTC CGG GTC AGG ACC 1125 1125 1134 1143 1152 1152 CCC GAG CCT GGG AAC CCT GTG GTG CAC TCG CCT GTT CAG GAT GAA CAC ACT GCC CCA CTC TGA 1191 1200 1209 1218 1227 1236 1245 GCC CTG CCG CTC GGC CTG GCA CCC TCG AAG CTA TGC GCC TAC GCG CTC AAG ATC TGG GCT CAC 927 936 945 954 954 963 GCC CTG GAG CGC GCT TTC GCG CTC TAC AAC CTG CTG GCC CTA TAC CTG CTG CTG CTC GCC ACC ACC ACC 663 672 ...681 690 699 708 717 GAT GGC GCC CTG CAG GGG CAG CTG CTA GCA CAG GGC GCT GGA GCA GTG CGC ACC AAG GTC TCC CGG 795 804 813 822 831 840 849 **16C GCC**

						FIG.3	ICRH	SVDR	YNLL	GPIQ	PHAS	FIC A
09	120	180	240	300	360	398	SLVIFV	ATLTAM	ERAFAL	LFAACW	PORORRI	
ALMLLGLVGN	GDFIACKFVNY	AAVSAPVLAL	LGRVAVRPAP	AGSWHPRSYA	PGPSDPAAPH		ALMLLGLVGN	IQQVSVQATC	CSEAFPSRAL	RH LGRAAVRPAP TDGALQGQLL AQRAGAVRTK VSRLVAAVVL LFAACWGPIQ	QAFCRVCPCG	
DAWLVPLFFA	LLYPLPGWVL	AVSLSIWVGS	CACYAAMLRH	LFLVLQALGP	APRRPRRPRR		DAWLVPLFFA	GDFMCKFVNY	HRLSPGPHTY	AQRAGAVRTK	LYAFLGSHFR	HTADI
DGPVPSPRAV	FLLCCVPFTA	LHRRTPRLAL	ALYLLPLLAT	LFAACWGPIQ	RQAFRRVCPC	LGEDNAPL	DGPGSAPRPL	LLYPLPTWVL	AAVSAPVLAL	TDCALQCQLL	SYSNSALNPL	VALCDCVODE
GCPGCGANAS	IANLAATDVT	WYYTVFPLRA	ERAFALYNLL	VSRLVAAVVL	LLYAFLGSHF	SGLAARGLCV	GCPGCGVNAS	FLLCCVPFTA	TVSLSIWVGS	LGRAAVRPAP	YALKIWAHCM	PATPED CND
ASWGAPANAS	SLVIYVICRH KPMRTVINFY IANLAATDVT FLLCCVPFTA LLYPLPGWVL GDFMCKFVNY	IQQVSVQATC ATLTAMSVDR WYVTVFPLRA LHRRTPRLAL AVSLSIWVGS AAVSAPVLAL	HRLSPGPRAY CSEAFPSRAL ERAFALYNLL ALYLLPLLAT CACYAAMLRH LGRVAVRPAP	ADSALGGQVL AERAGAVRAK VSRLVAAVVL LFAACWGPIQ LFLVLQALGP AGSWHPRSYA	AYALKTWAHC MSYSNSALNP LLYAFLGSHF RQAFRRVCPC APRRPRRPRR PGPSDPAAPH	AELLRLGSHP APARAQKPGS SGLAARGLCV LGEDNAPL	MAAEATLGPN VSWWAPSNAS GCPGCGVNAS DGPGSAPRPL DAWLVPLFFA ALMLLGLVGN SLVIFVICRH	KHMQTVTNFY IANLAATDVT FLLCCVPFTA LLYPLPTWYL GDFMCKFVNY IQQVSVQATC ATLTAMSVDR	WYYTVFPLRA LHRRTPRLAL TVSLSIWVGS AAVSAPVLAL HRLSPGPHTY CSEAFPSRAL ERAFALYNLL	CACYGAMLRH	LFLVLQALPL GGLAPSKLCA YALKIWAHCM SYSNSALNPL LYAFLGSHFR QAFCRVCPCG PQRQRRPHAS	AHSDRAABHS VEHSBAAHEV BVRIBEBSND VAHSBSVONE HIAPI
MHTVATSGPN ASWGAPANAS GCPGCGANAS DGPVPSPRAV DAWLVPLFFA ALMLLGLVGN	SLVIYVICRH	IQQVSVQATC	HRLSPGPRAY	ADSALQGQVL	AYALKTWAHC	AELLRLGSHP	MAAEATLGPN	KHMOTVTNFY	WYYTVFPLRA	ALYLLPLLAT CACYGAML	LFLVLQALPL	AHCUBAADHC

Leu CTG

75 75

AC AC AC

Leu CTG

క్టి

Tyr TAT 369

Ala Leu Leu 1 GCG CTC CTC 7 360

Phe Thr TTC ACC

Pro 851 351

Val GTA

Cys 760

Leu CTG

Le CH

Thr ACT

Val

Phe 110 333

Cys 760 342

Pro CCC 378

Val GTG 387

7/15

Leu CTG ပ္ပ Cys TGC Asp GAT Val Thr ACA 321 Phe TTC 255 구 55 8 GLy GGA 123 900 800 800 800 I le ATC A GCC Ser 100 A SS Val GTC Asp GAT Ala Leu / CTG (312 Leu CTG 180 Leu CTG 246 Asn AAC 114 Ite Ata Asn L ATC GCT AAC C 303 Ser TCA Ser TCC Gly Asn GGG AAC 237 Arg AGG Alg GCT 105 Pro CCA 171 Gly Leu Val (GGG CTA GTC (228 7,24 Ala GCG <u>다</u> Phe TC 7. 55 Ser TCC Asn AAT 294 Ser AGC 96 GLy GGC 162 a Ala Leu Met Leu Leu G T GCC CTA ATG TTG CTG G 219 AC S Gly Pro Val GTG Val GTC Asn AAC Asp GAT 153 Thr ACC 285 Pr 87 87 ca Gara Ser TCG GI, His Met (CAC ATG (276 Ala GCC Leu 176 Ala GCT 210 ACC ACC Asn AAT 144 His Lys CAC AAG Al GGG Val GTC Phe TTC Phe I GIY GGT Leu CTG 201 Arg CCC 267 Ala GCA 69 Cys 135 Cys 760 A GCC 5 5 5 5 <u>></u>၁ I le ATC Val GTG Met ATG 58

Th. A10 GCC 453 Cys TGT ACA ACA \$65 \$65 \$4 <u>-</u>58 Val GTG Ser 106 435 Val GTC GIn (CAG (426 Tyr Ile (TAC ATC Asn AAC 417 Val GTC Phe TTC Lys AA 408 Cys TGC Met ATG Phe TTC 399

FIG.5A

Arg CCC	2 88	Arg CGT	Thr	Asp GAT	Leu CTG	GIn	
					Arg CCC 849		
	Ser TCC	Pro CCC	Leu	Pro CCC	Ser TCC	Val GTG	
Leu	Val GTT	Phe TTT	Leu CTG	Ald GCA	Val GTC	Leu	
Ala GCA 510	Ala GCT 576	A10 GCG 642	Pro CCC 708	Pro CCC 774	AAG 840	Phe TTC 906	
Arg CCT	Ala GCA	G I u	Leu CTG	Arg CCC	Arg Thr 1	Leu CTG	
Leu CTG	Ser TCC	Ser AGT	Leu CTG	Val GTA	Arg 060	GIn	
582	÷156	\$ 50 53.33	75 S	÷ 25.89	515 53.50 53.50	11 e	
Phe TTC	Val GTG	Tyr	Leu	Ala CCC	g Alg Gly Alg V C GCT GGA GCA G 822	င် လ	
Val GTG	7.7 7.66	Thr	Ald	Arg CCC	GGA GGA	61y 660	
Thr ACT 492	11e ATC 558	His CAC 624	Leu CTG 690	61y 660 756	Ala GCT 822	77 766 888	4
Va GT	Ser	£ 53	Z E	Cle	₹8	Cys TGC	FIG 5B
Tyr TAC	Leu	61.y 666	Asn	His CAC	GIn CAG	Ala GCC	H
Trp 1GG 483	Ser AGC 549	Pro CCC 615	Tyr TAC 681	Arg CCC 747	Ala CCA 813	Ala GCC 879	
Arg CGC	Val GTC	Ser TCC	Leu CTC	Leu CTG	CTA	Phe TTC	
Asp GAC	Thr ACT	Leu CTG	₩ 88	Met ATG	Leu CTG	Leu CTC	
Val GTG 474	Leu CTG 540	Arg CGC 606	Phe TTC 672	Ala GCC 738	GIn CAG 804	Leu CTG 870	
Ser AGT	A la	His CAC	Ald	G1y GGT	61 <u>y</u>	Val GTC	
Met ATG	Leu CTG	Leu CTG	Arg CGC	Tyr TAC	GIn	Va! GTC	
Ala GCC 465	Arg CCC 531	Ala GCT 597	Glu GAG 663	Cys TGC 729	Leu CTG 795	Ala GCT 861	
Thr ACA	င် သ	Leu CTG	Leu CTG	A CC	A GCC	A CCC	
Leu TTG	Thr	Va I GTG	Ala	Cys 160	60 90 90 90	Val GTG	

H is CAC	Arg AGA	His	Thr	TGA
Alg GCT 981	She TTC 047	A1a 306 113	Arg AGG 179	Leu CTC 245
58	Ais I	Ser Ala TCA GCG 1113	Ad /	Pro Leu CCA CTC 1245
ATC	y Ser His Phe / T TCC CAC TTC / 8	DIA	7.9 20 20 20 20 20 20 20 20 20 20 20 20 20	Ala GCC
.ys 1	38 38	His /	o Val Arg Val Arg	Thr / ACT (
10 L	.eu ()	58	Pro V	lis 1
a Tyr Ala Leu Lys C TAC GCG CTC AAG / 963 972	Phe Leu Gly TTC CTG GGT 1038	7 9 50 00 0	His F	GAA CAC
ут АС В 63.	20 E	7.79 A 67 C 95 C	-88- -88-	sp GAT GAT G
- L & - S	1 Tyr Alg 2 TAT GCC 1029	Gly Pro Gln Arg Gln Arg Arg Pro His GCC CCC CAA CCC CAG CCT CCG CCC CAC 1086 1095 1104	s Ser Arg Ala Ala C AGC CGG GCT GCG 1152	o Ser Val Gln Asp G C TCT GTT CAG GAT G 1218
χ SC S S	TC T	<u>කි</u> රි ල උ	ნე ტე	19 E
r Lys Leu Cys Ala G AAG CTA TGC GCC 954	Pro Leu Leu CCC CTC CTC 1020	- A - B - B - B	er A GC C 52	er V CT G 18
က် သည်	58 	5 လ ၁ ၁ ၁	His S CAC A	20 S 12 12
≨ک	£ 5	<u> </u>	主の	P C C C C
Ser TCS	ASI AA(61y 660	58 58	Ser TCC
A CCC TCC 945	Leu CTC 1011	Cys 160 1077	Val GTG 1143	His CAC 1209
₹33	r Ala Leu T GCG CTC 7	8 Pro Cys CCC TGC 1077	Ser AGT	Val GTG
Leu CTG	Ser	Cys TGC	His CAT	Val GTG
Gly Leu GCC CTG 936	Asn AAT 1002	Val GTG 1068	r CCC CAT AGT GTG (1134	CCT GTG GTG CAC 1200
<u>></u> 99	Ser	Arg GC	Ala GCA	Asn AAC
Leu CTC	Tyr TAC	Cys TGC	A GCC	GE 9
Pro Leu (CCG CTC (927	Ser 1CC 993	Phe TTC 1059	Arg CGA 125	Pro CCT 191
Leu CTG	Met ATG	A CCC	Asp Arg GAC CGA 1125	Glu Pro GAG CCT 1191
Ala Leu I GCC CTG	Cys TGC	GIn	Ser TCC	Pro CCC
•				

FIG.5C

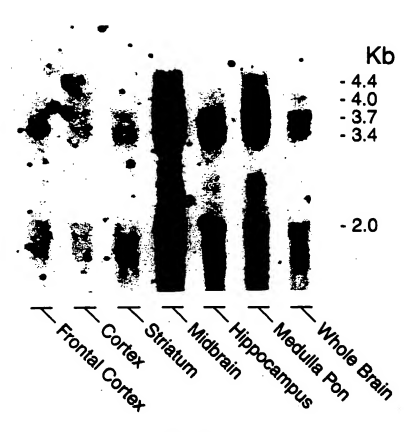


FIG.6

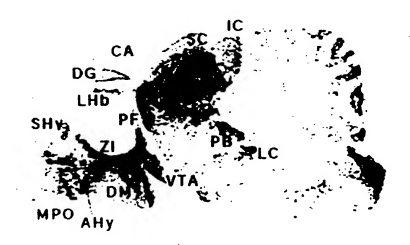


FIG.7A

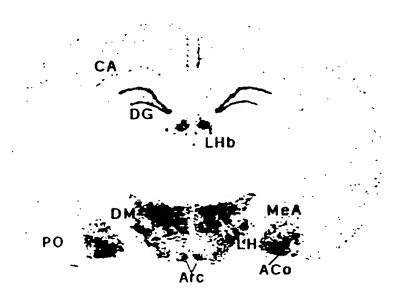


FIG.7B

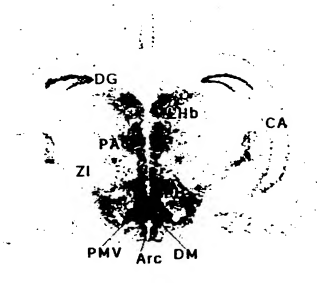


FIG.7C

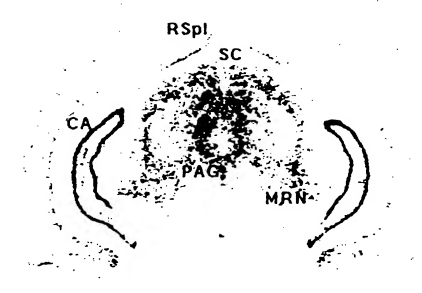


FIG.7D

SUBSTITUTE SHEET (RULE 26)

33 23 16 8 39	116 108 99 96 122	200 194 183 180 209	288 268 259 258
MAREATLGPNVSWWAPSNA-SGGNGCGVFASIDGP- MELAPWNL-SEGNGSDPEPPAEPR MNGSGSQGAENTSQEG	TRANSMEMBRANE 1 -GSAFRPL DAME VPLEFAAL MELGE VGNSE VIFVICP KAMOTIVTN-FYTANLAATON TELLCCVFFTAFLNTAL PTWVLGDFMCK -PLFGIGVENFITL VYFGL IF AMGVLGNSL VITTN-APSKPGKPRSTTNLF-TILINLS I ADLAYLLECIFFGAT VYAL PTWVLGAFICK -GSIGGVOPEANL VPLFFAL IFLUGAVGNAL VLAVLLOGGSAVGEPRSTTNLF-TILINLGVADLCFTILCCVFFGAT INTILIDINYFIGSLUCK -LDSFGSVGAVAVPVIFAL IFLLGAVGNAL VLAVLLOGGPSAVGEPRSTTNDLF-TILINLAVADLCFTILCCVFFGAT INTILIDIANLFGEVCK ARSASSLALAJIA I TALYSAVCAVGLLGNAL VVEGTINPYTKLKTIATN-IYIFNLALAD-ALATSILFFDSAKYLMETWPFGELLCK	TRANSMEMBRANE 3 FUNNTIOCOVSVORTCATL TAMSVDRWYVTWFPLRALHRRTPRALIT-VSLSIWVGSAAVSAPVLALHRLSPGPHTY-CSEAFPSRAL FINNTIOCOVSVORTCATL TAMSVDRWYVTWFPLRALHRRTPRALL GWGF-IMALSIJAMASPVAYYORLFHRDSNOTI-FOWEHWPNOLH FINNTIOCOVSVORLFTLAAMSVDRYVATWFLHSRELRIPRNALAAIGC-IMGLALLFSGPYISYYRDSOL-ANLIT-VCHPAWSAPR AWHFLIYLTMHASSFTLAAAWSLDRYLAVRHOLRSRALRIPPNARAAWGL-IMGLALLFSGPYLSYYRTSOL-ANLIT-VCHPAWEDAR AWHFLIYLTMYASSFTLAAAWSLDRYLAVRHOLRSRALRIPPNARAAWGL-YWLLAALFSGPYLSYYGTVRY-GALEL-CVPAWEDAR AWLSIDYYNMFTSIFILTMSVDRYIAVCHPVKALDFRIPAKAKLINIC-IWVLASGWSVPIMWMAVIOPRDGAVVCTLOFPSPSWYW	TRANSMEMBRANE 5 ERAFALIVNILALITLIPLIATGACYGAMLRHLGRAAVPPAPTDGALQGQLLADRAGAVFTKVSPLIVAAVVILFAACVCPIJQLFLIVLIQALKKAYVVCTFVFGYLLPLULIGFCYGAMLRHLGRAAVPPAPTDGALQGGLLADRAGAVFTKVSPLIVAVVVLLFAACVCPIJQLFLIVLA-RRAMDLCTFVFGYLLPVUVLSUTYARTLRNLWR-TVV-DFVTAGSGSGRAKRKVIIFPMIIIJVAVLFCLCVWPHHA-LIILCV-RPPLDVATFAAGYLLPVAVVSLAYGRTLGFLWA-FAVGPA-GGAAAGEARPRATGFAGPAMLAVAALYALCVGPHHA-LIILCF-DTVTKICVFLFAFVVPIILIITVCYCLMULRLRSVPLLSGSKEKDRSLRRITPMILVVVGAFVVCWAPIHIEVIVV
iPRS4 ia IR1 ia IR2 ia IR3)OR	JPR54 Ja 1R1 Ja 1R2 JDR	GPR54 Ga IR1 Ga IR2 Ga IR3 DOR	GPR54 Ga IR1 Ga IR2 Ga IR3 DOR

SUBSTITUTE SHEET (RULE 26)

TRANSMEMBRANE 7

377 346 348 347 372	395 372 370
PLGGLØPSKLCAYALKIJWAHOMSYSNSALNPLLVAFLGSHFROAFICRVCPCGPGRORPPHASAHSDPAA-PHSVPHSRAAHFVBVRTPFP EFIGAFPLTPAS-FFFRIJTAHGLAYSNSSVNPTITVAFLSENFRKAYKÖVFKCRVCNESPHGDAKEKNRTDTPFSTNCTHV VFIGRFPLTRA-TYALRIJLSHLVSYANSCVNPTVALVSKHFRKGFRKTCAGLLRPAPRRASGRVSJLAPGNHSGSMLEGESTDLTQVSEA VYGRFAFFSPA-TYACRLASHGLAYANSCLNPLVVSLASRHFRARTRRLVPCGRORHRHHFRAHRALRRVCPASSGPAGYPGDAPPRGVSM TLVDINRRDPLVVAALHLCTALGYANSSLNPVLYAFLDENFIKRCFROLCRAPCGGCFPGSLRRPRGATARERVTACTPSDGPGGAAA	4 GNPVVHSPSVQDEHTAPL 2 AGPLNPPPALPNCTASSRTLDPAC 3 EPRGDALRGGGETRLTLSPRGPQ
GPR54 Ga IR1 Ga IR2 Ga IR2 DDR	GPR54 Ga 1R2 Ga 1R3
CUE	CTITUITE C

FIG.8B

SUBSTITUTE SHEET (RULE 26)

WO 00/50563 PCT/US00/04416

15/15

rat	10 MAAEATLGPNV	20 SWWAPSNASGCF	30 PGCGVNASDGF	40 PGSAPRPLDAV	50 VLVPLFFAALI	60 NLLGLVGN
humon	: :: ::: MHTVATSGPNA 10	:: :::::: SWGAPANASGCF 20	CCGANASDGF 30	: :::::: PVPSPRAVDAV 40	::::::: VLVPLFFAAL! 50	MLLGLVGN 60
rat	70 SLVIFVICRHKI	:.:::	• • • • • • • • • • •		:::: :::::	:::::::
human	SLVIYVICRHKI 70	PMRTVTNFYIAN 80	NLAATDVTFLL 90	.CCVPFTALL) 100	PLPGWVLGDF 110	FMCKFVNY 120
rat	130 IQQVSVQATCA		150 /TVFPLRALHR		170 SLSIWVGSAAV	180 /SAPVLAL
human	IQQVSVQATCA 130	TLTAMSVDRWY\ 140	/TVFPLRALHR 150	RTPRLALAVS 160	SLSIWVGSAA\ 170	/SAPVLAL 180
rat	190 HRLSPGPHTYCS	200 SEAFPSRALERA	210 FALYNLLALY	220 LLPLLATCAC	230 CYGAMLRHLGR	240 RAAVRPAP
human	HRL SPGPRAYCS	SEAFPSRALERA 200	FALYNLLALY 210	LLPLLATCAC 220	YAAMLRHLGR 230	RVAVRPAP 240
rat	250 TDGALQGQLLAC		270 LVAAVVLLFA			LAPSKLC
human	ADSALQGQVLAE 250	RAGAVRAKVSR 260	LVAAVVLLFA 270	ACWGPIQLFL 280	VLQALGPAGS 290	: WHPRSYA
rot	300 310 AYALKIWAHCMS	320 YSNSALNPLLY	330 AFLGSHFRQA	340 FCRVCPCGPQ	350 RORRPHASAH	SDRAAPH
human	AYALKTWAHCMS 310		'AFLGSHFRQA 330		RPRRPRPGP 350	SDPAAPH 360
rat	360 370 SVPHSRAAHPVR	380 VRTPEPGNP—	390 VVHSPSVQDE			•
humon	AELLRLGSHPAP 370	ARAQKPGSSGL 380	AARGLCVLGE 390	DNAPL		

FIG.9

INTERNATIONAL SEARCH REPORT

International application No. PCT/US00/04416

IPC(7)	SSIFICATION OF SUBJECT MATTER C12N 5/10, 15/00, 15/09, 15/11, 15/12, 15/63 435/320.1, 325; 536/23.5				
	o International Patent Classification (IPC) or to both n	ational classification and IPC			
B. FIEL	DS SEARCHED				
Minimum d	ocumentation searched (classification system followed	by classification symbols)			
U.S. :	435/320.1, 325; 536/23.5				
Documentat	ion searched other than minimum documentation to the	extent that such documents are included	in the fields searched		
Electronic d	ata base consulted during the international search (na	ne of data base and, where practicable,	search terms used)		
Please Sec	e Extra Sheet.				
c. Doc	UMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where app	ropriate, of the relevant passages	Relevant to claim No.		
X	WO 98/03548 A1 (ASTRA PHARM	A, INC.) 29 January 1998	3		
-	(29.01.98), especially pages 8-9.				
Α			1-2, 4-5		
X, P					
	receptors," FEBS Letters. 05 March 1 103-107, especially page 103 and figur				
	103-107, especially page 103 and figur	cs 1-3.	,		
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Furtl	ner documents are listed in the continuation of Box C.				
i e	pocial categories of cited documents:	"T" later document published after the inte date and not in conflict with the appl	ication but cited to understand		
	ocument defining the general state of the art which is not considered be of particular relevance	the principle or theory underlying the			
	rlier document published on or after the international filing date	"X" document of particular relevance; the considered novel or cannot be considered.	red to involve an inventive step		
C1	ocument which may throw doubts on priority claim(s) or which is ted to establish the publication date of another citation or other	when the document is taken alone "Y" document of particular relevance; the	claimed invention cannot be		
•O• do	special reason (as specified) considered to involve an inventive step when the document is				
	ocument published prior to the international filing date but later than a priority date claimed	*&* document member of the same patent	family		
Date of the	actual completion of the international search	Date of mailing of the international sea	rch report		
15 AUGU	JST 2000 .	07 SEP 2000			
Name and	mailing address of the ISA/US	Authorized officer my Mar	2		
Commission Box PCT	oner of Patents and Trademarks	MIZHAEL PAK			
	on, D.C. 20231 No. (703) 305-3230	Telephone No. (703) 308-0196			

INTERNATIONAL SEARCH REPORT

International application No. PCT/US00/04416

Box	ιo	bservations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This	interr	national report has not been established in respect of certain claims under Article 17(2Xa) for the following reasons:
1. [Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. [Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. [Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box	11 (Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This	Inter	mational Searching Authority found multiple inventions in this international application, as follows:
	Ple	ease See Extra Sheet.
		·
1,		As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2.		As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.		As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4.	X 1	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Ren	nark	on Protest
		No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

International application No. PCT/US00/04416

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

BRS, STN, MEDLINE, GENBANK, PIR. EMBL, GENESEQ, EST

search terms: g-protein receptor, galanin receptor, cDNA, GPCR, GPR54

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s) 1-5, drawn to an isolated DNA, expression vector, and recombinant host cell.

Group II, claim(s) 6-9, drawn to an isolated polypeptide.

Group III, claim(s) 10-11, drawn to a method of determining whether a substance is an agonist or antagonist by binding. Group IV, claim(s) 12, drawn to a drawn to a method of determining whether a substance is an agonist or antagonist by functional response.

Group V, claim(s) 13, drawn to an antibody.

and it considers that the International Application does not comply with the requirements of unity of invention (Rules 13.1, 13.2 and 13.3) for the reasons indicated below:

The inventions listed as Groups I-V do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features because the product of claim 3 is anticipated by ASTRA PHARMA INC. (WO 98/03548. 29 January 1998. pages 7-8) and thus, does not share a special technical feature with any other group.

The special technical feature of Group I is a DNA, an expression vector, and a host cell. Pursuant to 37 CFR 1.475(d), these claims are considered by the ISA/US to constitute the main invention, and none of the related groups II-V correspond to the main invention.

The products of Group II and V do not share the same or corresponding special technical feature with Group I, because they are drawn to products having materially different structures and functions, and each defines a separate invention over the art.

The methods of Groups III and IV, do not share the same or corresponding special technical feature with Group I, because the methods have materially different process steps and are practiced for materially different purposes, and each defines a separate invention over the art.

Since Groups I-V do not share a special technical feature, unity of invention is lacking.